COLLEGE COLLEGE CHEMISTRY

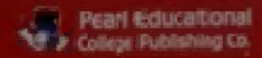
Intermediate Part I





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College Chemistrus Part 1

Chapter-1

Basic Concepts

ATOM

Evidence for an atom Molecule lon

Molecular lon

RELATIVE ATOMIC MASS ISOTOPES

Relative abundance of laotopes

Mass spectrometry Fractional atomic masses

ANALYSIS OF A COMPOUND

Empirical formula Empirical formula from combustion analysis Molecular formula

MOLE

Avagadro's number Molar volume

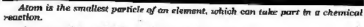
STOICHIOMETRY LIMITING REACTANT

Determination of limiting reactant

YIELD.

Objective and short answer, quartions (exercise) Numerical problems (exercise) Past Papers MCQs and Short Questions

Test your skills



- The atoms of He, Ne (noble gases) can exist independently.
- The atoms of $\underline{\text{hydrogen}}$, nitrogen and oxygen cannot exist independently.
- Atom is now further divisible into sub-atomic particles.
- More than 100 sub-atomic particles are thought to exist in an atom.
- A few sub-atomic particles are electron, proton, neutron, hypron, neutrino, antineutrino
- Electrons, protons and neutrons are the fundamental particles of atom.

History of Alom

Earlier Concept and Greek Philosopher's Work

- Earlier it was though that matter is made up of simple, indivisible particles.
- Greek philosophers thought that matter could be divided into smaller and smaller particles. Thus a basic unit could be reached which cannot be further sub-divided.
- Democritus (460-370 B.C.) called the indivisible particles of matter as atoms
- The word atom is derived form Greek word 'atomos' means 'indivisible'.
- The concepts of Greek Philosophers were not based on experimental proofs.

Late 17th Century work

- In late 17th century, the quantitative composition of substances was studied, it showed that few elements are present in many different substances.
- It was also studied that how elements combine to form compounds and compounds could be broken into elements.

Daltan's Work

- In 1808, John Delton, an English School teacher, explained law of conservation of mass and law of definite proportion on the basis of concept of atoms. He developed an atomic theory.
- According to main postulate of <u>Politon's theory</u>, all matter is composed of atoms of different elements, which differ in their properties.

Berselius Work

- A Swedish chemist J. Berzellus (1779 1848) determined the atomic masses of elements. A number of his values are very close to modern values of atomic masses
- Berzellus developed the system of giving symbols to elements.

Evidence for an Atom

Use of Optical Microscope:

Ordinary optical microscope can measure the size of an object up to or above 500 nm. Therefore, a clear and accurate image of object that is smaller than the wavelength of visible light cannot be obtained. Since atoms are much smaller than 500 nm, So, they cannot be seen with optical microscope.

Use of Electron Microscope:

A direct evidence of atom can be obtained by an electron microscope. If uses a beam of electrons instead of visible light. The. wavelength of electrons is much shorter than visible light and is suitable to see extremely small objects. A less clear picture of atoms is obtained by an electron microscope.

Example: The figure shows an electron microscopic photograph of a piece of graphite. It has been magnified about 15 million times. The bright bands in the figure are layers of carbon atoms.



Use of X-reper

in the 20th century, X-ray work has shown that

(f) the <u>diameters</u> of atoms are approximately 2×10^{-18} m or 0.2×10^{-9} m or 0.2 rim.

- The masses of stoms are often expressed in atomic mass units (amu).
 - $1 \text{ amu} = 1.661 \times 10^{-87} \text{ kg} = 1.661 \times 10^{-87} \text{ g}$
- A full stop may have two million atoms present in it.

MOLLCOLE

The smallest particle of a pure substance, which can exist independently, is called a molecule.

Arosinto (Monatennic and Palentunic molecules)

- The number of atoms in a molecule is called its atomicity.
- A molecule containing one atom is called a monostomic molecule,
- A molecule containing two or more atoms is called a polystomic molecule. The polyetomic molecules containing two or three atoms are called diatomic and triatoms: molecular respectively.



College Chamberry, Port-J

Examples:

- Monatomic molecules: He, Ne, other noble gases
- Polyatomic molecules: H₂, CO, O₁, CO₂, NH₃ etc.

Homosatomic and Heterisatomic indecates

(Difference between molecule of an element and compound)

- Molecules of elements may contain one, two and three same type of atoms, e.g. He, Cla. O₃, P₄, S₆. These are called homo-atomic molecules.
- Majecules of compounds consist of different kind of atoms. e.g. HCl, NH₅, H₆SO₆, CoH11Os. These are called hetero-atomic molecules.

Microundeedes and Macronal cutes

- Molecules have bigger size than atoms. The size of molecule depends upon number of atoms in a molecule. (N) shape of molecule.
- Molecules containing fewer numbers of atoms and are small in size are called micromolecules.

Examples: CO₂, NH₂ H₂SO₄, C₆H₁₃O₆ etc.

Molecules containing large number of atoms and are large in size are called macromolecules.

Example: Haemoglobin: It is found in blood. It helps to carry oxygen from lungs to all parts of body. One molecule of haemoglobin is made up of about 10,000 atoms, it is 68,000 times heavier than a hudrogen atom.

ions are those species which carry either positive or negative charge.

These are of two types

Paratire fairs

Ione which carry positive charge are called positive ione.

When an atom of an element loses one or more electrons, positive ions are produced.

A* + &

The removal of electrons from an atom requires energy, hence formation of positive ion is an endothermic process.

Positive lone are called cations.

- An atom may loss one, two or three electrons to form monopositive, dipositive or impositive lone respectively.
- The most common positive ions are formed by the metal atoms

Emmules:

Na" , K", Ca** , Aft*, Fe1*, Sn4* etc.

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Beste Concess

Segundo dessi-

ions which carry negative charge are natled negative lone.

 When an atom of an element picks up one or more electrons, negative ions be produced.

B + # --- B

- The addition of one electron in an atom usually results in release of energy. It is due to
 the attraction between nucleus and incoming electron. Hence formation of uninequity
 ion is an exothermic process.
- An atom may gain two electrons to form a dinestative fort. However, it will be an
 endothermic process due to reputation between uninequative ion and second incoming
 electron.
- Negative ions are called Anions.

Examples

HOTE:

- Negative ions may constat of group of atoms e.g. CO₃¹*, Cr₆O₂¹*, MnO₄*, PO₄³*, SO₄³* etc.
- Positive tons containing group of atoms are rare e.g. NH₄*, H₈O* and carbocations in organic chemistry etc.
- The cations and anions have completely different properties from their corresponding hearts access

West order los

The low fermed due to low or gain of electrons by a molecule is called a molecular lot Examples:

CH₄*, CO*, N₄* etc.

Abundance: Cotionic molecular ions are more abundant than anionic molecular ions.

Molecular ions are generated by passing high enemy electron beauti schemicles or X-rays through a gas.

Application: Breaking of molecular ions from natural product can give important information about structure of the compound.

PLIANVE ALOME MASS

Definition: It is the mass of an atom of an element as composed to the mass of an atom surbon taken as 12.

Unit: Alomic mass is expressed in ATOMIC MASS (BITTS, denoted by amu.

Culture Chamberry Eggs

The $\frac{1}{12}$ of the more of one atom of curbon is collect 1 and

1 amu = 1.661 × 10⁻²⁷ kg = 1.661 × 10⁻⁸⁶ g

Examples:

Atomic mass unit is defined as

On carbon-12 scale, the relative etomic mass of $^{12}_{\circ}$ C is 12,0000 arms and that of |H| in 1,0078 arms.

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Why relative atomic mass unit scale is used?

Atomic masses of atoms are very <u>small</u>. These <u>cannot</u> be measured by a <u>phosical</u> balance. Thus relative atomic mass unit scale is used.

Lable Didative Atomic Masses of a few elements Relative Relative						
Liennam	Alumn Shass	Homeots	Atomic Mas			
н	1.00794	a	35.453			
O.	15.9994	Cu	63.546			
Na	20.1797	U	238.0289			

Partory!

John Dahon thought that all the atoms of an element were similar. However, Soddy showed that alons of the same element may be different.

Dafinition: Atoms of the same element, buting same atomic number but different stants masses are called isotopes

The phenomenon of isotopy was discovered by Soddy.

Properties of Isotopes

- isotopes have same number of electrons and protons but different number of peutrons in their nuclei
- ⁴ linitopes of an element have <u>different physical properties</u> due to different number of neutrons in their nuclei.
- a isotopes of an element have <u>same chemical properties because they have</u> seme number of valence electrons. During chemical reactions valence electrons are involved.
- * Isotopes have same position in the periodic table since they have same stornic number.

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to Carbon has three tectopes written as \$10. \$10. \$10. and expressed as C-12, C-13, C-14. having 6, 7 and 8 neutrons in their nuclei respectively.

(a) Hydrogen has three isotopes

H Produm

H Deutertum H Tritium

Occurry has three inotopes, Nickle has first isotopes, Calcium has six isotopes, Pallodium has sig incropes. Codmitten has nine isotopes, tip has eleven isotopes etc.

Schothe Alamahan e of Lastopes

- The natural occurrence of different isotopes of an element is called their relative abundance, botopes of all the elements have their own natural abundance.
- The masses and abundance of isotopes of an element can be determined by Mass Spectrometry.
- The properties of elements are mostly like that of most abundant isotope of that element.

Facts about Eduting abundance of local

- There are above 280 different isotopes that occur in nature.
- Out of 280, there are about 40 radioactive isotopes
- Buildes these, about 300 unstable radioactive isotopes have been produced through artificial disinfegration in nuclear reactor.
- Elements with <u>odd atomic number</u> almost have maximum <u>two</u> stable isotopes
- Elements with even atomic number usually have many isotopes
- Isotopes with mass number of four or some multiple of four are more abundant e.g. 16O, 16Mg, 16Si, 16Ca, 16Fe form nearly 50% of the earth crust.
- Out of 290 natural innicone , 154 have gven mass number and gven atomic number

Stangerstager Diameters

y only ant lectors are called mono-isotopic elements.

Amenic, Pluorine, lodine, Gold etc have only single isotope.

e Chamleton Part-I

Fable Namual abundance of some common isotopes					
## - # Elements	Isotope	Abttechate to ("a)			
Hydrogen	'H, "H	99.985, 0.015			
Carbon	12C, 12C	98.893, 1.107			
Nitrogen	MM MM	99 634, 0.366			
Oxygen	™O, ™O, ™O 99.759, 0.033				
Sulphur	#25, #85, MS, MS	95.0, 0.76, 4.22, 0.014			
Chlorine	Mai na	75.53, 24.47			
Bromine	Migr. at Br	50.54, 49.49			

MASS SPECIROMETRY

Mass Spectrometry is an analytical method used to measure the exact masses different isotopes of an element

The instrument used for this purpose is called Mass Spectrometer

- Initially Aston's mass spectrograph was used to identify isotopes on the basis of their alomic masses
- Dempster's mass spectrometer was designed for the elements found in solid state.

Method

A typical mass spectrometer consists of following parts

(1) Vaportzaiton Chamber

(2) Ionization Chamber

(3) Analyzer

(4) Electrometer (Ican Collector) (5) Detector, Amplifier and Recorder

(I) Vaportsation Chamber

The substance is first converted into vapour stats. The pressure of these vapours is kept vary low i.e. 10^{-6} to 10^{-7} torn.

(2) Ionization Chamber

The vapours are then passed into an ionization chamber. In this fast movine electrons are thrown upon them. As a result gaseous atoms are ionized and positive ions are produced. These positive ions have different masses due to presence of different isotopes in them.

(2) Analyzer

The analyzer separate ions on the basis of their m/e value in two steps.

(i) Acceleration of loss: Each positive ion has its own mic (mass to charge) value. A potential difference (E) of 500 - 2000 volts is applied between the perforated plates to accelerate the ions. Thus ions are strongly attracted towards the negative plate.

(a) Deflection of Issue: The ions are then pussed through a strong magnetic field of strength "If It defects the ions on the basis of their mire value. Thus, jons follow a circular path,

The mathematical relationship is

$$m/e = \frac{H^2r}{E}$$

Where H = Strength of magnetic field,

- E =Swength of electric field,
- r = radius of circular path.





By heeping 'E' constant, if 'H' is increased, 'Y' also increases. Thus, lons of a particular m/e value can be made to fall at different place than before.

- It can also be done by keeping "H" constant and by changing "E".
- So, smaller the 'm/e' value, smaller will be the 'r' produced by magnetic field. Hence ions with same rate values are separated into beams of ions.

(4) Electrometer (Ion Collector)

In mass spectrometer, the separated beams of ions are passed through allt one by one Each beam contains ions of specific m/e value. These fall on an electrometer, called ion collector. The electrometer develope electric current. The strength of current for each beam gives the relative abundance of ions.

(E) Detector. Amplifier and Recorder

in modern spectrographs, the ions strike the detector to generate current. The current is ampitfied and led to the computerized recorder.

19 Comparison with C-12: The same experiment is performed with C-12 isotope and the current strength is compared. This comparison

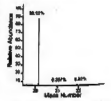
gives the exact mass number of isotope.

Exercise Q8 (b): Flow does a mass spectrograph show the relative abundance of isotopes of an element?

(II) Mase Spectrum (or Mose Spectrograph):

In mass spectrometer, the result is recorded in the form of a graph containing peals. The relative numbers of lone are present along V-axis (ordinate) and m/e values of ions are present along X-axis (absciasa). This is called Mass Spectrum, it gives following information

- the number of peaks gives the number of isotopes of an glement
- the height of peak gives the teletive abundance of isotopes
- the position of peak gives the mass number of isotopes.



Fig(1.3) Computer plotted grips for the leatonsh of above

1.1

Other techniques for superetion of tease

isotopes can also be separated by following techniques on the basis of their properties. (r) gaseous diffusion (n) thermal diffusion (III) distillation.

(to) ultracentrifuge

(v) electromagnetic suparation and

(at) laser separation etc.

desic Principle of Mass Spectrometre

A substance is first volatilized. It is then ionized with the help of high energy beam of electrons to form gaseous positive ions. These ions are separated on the basis of their mass to charge ratio (m/e). The result is recorded in the form of peaks. The result is a graph in which mile is plotted as abscissa (x-axis) and the relative number of ions as ordinate (v-axis). This graph is called mass spectrum. It tells about number, mass and relative abundance of BOTODES.

Inetional Atomic Masses

Exercise Q5.(a)

What are the isotopes? How do you deduce the fractional atomic masses of elements from the relative isotopic abundances? Give two examples in the support of your answer?

The alumic masses of elements are written as average atomic masses. These depend upon the number of isotopes of an element, their masses and their natural abundance. Thus, most of the elements have fractional atomic masses.

A sample of moon to found to constat of 10 Ne, 10 Ne and 10 Ne in the percentages of 90.92%, 0.26%, 8.82% respectively.

The average atomic mass of Ne is given as

Average atomic mass =
$$\frac{(20 \times 90.92) + (21 \times 0.26) + (22 \times 8.82)}{100} = 20.18$$

Hence the average atomic mass of neon is 20:18 amu

Actually, there is no individual neon atom in the sample that has a mass of 20.18 arms. However, in mutine work the sample of neon is considered to have average mans of 20,18 a.m.u.

Situar has too important naturally occurring leatopus, ¹⁶⁷Ag and ¹⁶⁹Ag with relative abundances 51.84% and 48.16% respectively.

The average atomic mass of Ag is given on

everage atomic mass =
$$\frac{(107 \times 51.84) + (109 \times 49.16)}{100} = 107.96$$
 amu

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Book Compley

ANALYSIS OF A COMPOUND

To find the molecular formula of a compound following steps are taken.

- 1. All the elements present in the compound are identified. This is called qualitative analysis.
- The mass of each element is determined in the compound. This is called quantilable analysis.
- The mass of each element is used to calculate the <u>percentage by mass</u> of each element.
- 4. The percentage is used to determine the empirical formula of the compound
- 5. Finally molecular formula is obtained from empirical formula and molecular may.

Persontage of Element in a Companied

It is the number of grams of an element present in 100g of the compound.

 Percentage from given amounts: The percentage of each element in a compound as be determined by following formula.

$$Percentage of an element = \frac{Mass of the element}{Mass of the compound} \times 100$$

Percentage from Formula Mass: The percentage of each element in a compound can't
determined theoretically from formula mass of a compound.

Example 2:

8.657 g of a compoundy were decomposed (rito its element and gave 5.217 g of carbo-0.962g of hydrogen, 2.478 g of oxygen. Coldulate the percentage composition of the compound under study.

Solution:

→ Mass of the given compound = 8.657g

Mass of C = 5.217g

Mass of H = 0.962g

Mass of O = 2.478g

Parcentage of the elements can be calculated by the formula.

% of element = \frac{\text{mass of the element}}{\text{mass of the compound}} \times 100

College Chemistry: Pert-)

Books Concepts

% of C =
$$\frac{5.217}{8.657} \times 100 = 60.26\%$$

% of H = $\frac{0.962}{0.228} \times 100 \times 11.11\%$

% of
$$O = \frac{2.478}{8.657} \times 100 = 28.62\%$$

8.657

¢

Empirical Formula

The formula, which shows the simplest whole number ratio between atoms of a compound, is called sempirical formula.

Examples:

Empirical formula of hydrogen peroxide is HÖ. It shows that simplest whole number ratio between H and O is 1:1.

Similarly empirical formulas of glucose and benzene are CH₂O and CH respectively.

Steps to Determine the Empirical Formula

- (i) Determine the percentage composition of each element in a substance
- (III) <u>Divide</u> the percentage of each element by its <u>atomic mass</u> to get <u>number of gram atoms (moles)</u>.
- (All) <u>Ovide</u> the moles of each element by the <u>smallest</u> number of moles to get <u>atomic</u> ratios.
- (Ne) If atomic ratios <u>are not</u> in simple <u>whole number</u>, then multiply with a small <u>suitable</u> <u>number</u> to get whole number ratio.

Thus empirical formula is obtained.

Jaleeular Formula

The formula, which shows the exact number of atoms of each element present in one majacule of a compound, is called molecular formula.

it is based on an actual molecule

Econopies:

Molecular formulas of bencene and glucose are C₆H₆& C₆H₁₈O₆ respectively.

R: I: tiouship Better on Empireal and Molecular Formula

 Molecular formula may be the multiple of empirical formula. For many compounds amplitude and molecular formulas are different.

Becompley: Empirical formulas of benzene and glucose are CH and CH₀O respectively. However, their molecular formulas are C_0H_0 and $C_0H_{12}O_0$ respectively.

This relationship can be expressed as

Molecular lumula = n (empirical formula)

where, 'e' is an integer and its value is 1, 2, 3,....

The 's' shows the ratio of molecular mass and empirical formule mass.

Molecular mass n - Empirical formula mass

(H) The compounds may have same empirical and molecular formulas. Brownplace NHs, HsO, COs, CrsHssOss etc. For such compounds value of 'n' is unity.

Mr.

n between Empirical and Molecular Formule

Distance parment Caparica	Modeenter Estimate
It shows the simplest whole number ratio between alone of a compound.	It shows exact number of atoms of each element present in one molecule of a compound.
Both ionic and covalent compounds have emprical formula	lorac compounds do not have molecular formula.
It is based on formula unit which may or may not exist independently	It is based on an actual molecule which exist independently
It is obtained from percentage composition of elements in a compound.	It is obtained from empirical formula by using the relationship. Molecular formula = n (empirical formula)
Engriples: Empirical formulae of glucose and benzene are CH ₂ O and CH remetively.	Examples: Molecular formulae of glucose and benzene are C ₆ H ₁₈ O ₆ and C ₆ H remortively.

EMPIRICAL FORMULA FROM COMBUSTION ANALYSIS

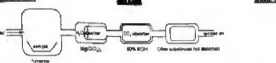
umbustion Applysis:

It is an experimental procedure by which amounts of various elements present in the semination of a compound are determined by burning.

Organic compounds containing only C. H and O are analyzed by combustion analysis. Method

- A weighed quantity of the compound is burned, in a combustion tube. The combustor tube is litted in a furnace.
- Oxygen is supplied to burn the compound.
- Hudrosen and carbon contained in the compound are converted to Hu2 & Q0

C & H in sample + O₂ → CO₂ + H₂O



- . In combustion analysis, the water vapours formed are absorbed in magnesium perchiorate [Mg(ClO.).
- The CO, is absorbed in 50% KOH solution.
- The increase in masses of these absorbers gives the masses of HaO and CO, produced.
- Other absorbers can be used for other products of combustion.
- The amount of oxugen is determined by the method of difference.
- Formulas for calculations: For a compound containing C. H & O only, the formulas for percentages are

% of C =
$$\frac{\text{mass of CO}_2}{\text{mass of organic compound}} \times \frac{12}{44} \times 100$$

% of H =
$$\frac{\text{mass of } H_2O}{\text{mass of organic compound}} \times \frac{2.016}{18} \times 100$$

% of oxygen = 100 - (% of carbon + % of hydrogen)

Ascerbic acid (ultanin C) contains 40.92% carbon, 4.58% hydrogen and 54.5% of Oxygen by mass. What is the empirical farmula of the oscorbic seid?

(Gujrenurski Board, 2010: Multan Board, 2007: Bahawaipur Board, 2011)

Sofetton:

Lichen		No. of presentations	Atomic Reto
С	40.92	$\frac{40.82}{12} = 3.41$	$\frac{3.41}{3.41} = 1$
н	4:58	$\frac{4.58}{1.008} = 4.54$	$\frac{4.54}{3.41} = 1.33$
0	54,50	$\frac{54.5}{16} = 3.41$	$\frac{3.41}{3.41} = 1$

Multiply atomic ratio by 3 to get values in simple whole numbers.

Thus the empirical formula for ascorbic acid is C.H.O.

all containing carbon, hydrogen and suggest only use subjected to also 0.5439 g of the compound grow 1.039 g of CO_p , 0.6369 g of H_2D . ile of the compound.

Mass of organic compound = 0.5439g

Mass of CO₂ = 1.039g

Mass of $H_1O = 0.6369g$

 $= \frac{1.039}{0.5439} \times \frac{12}{44} \times 100 = 52.10\%$ Thus Percentage of C

 $\frac{0.6369}{0.5439} \times \frac{2.016}{18} \times 100 = 13.12\%$ Percentage of H

Percentage of O = 100 - (52.11 + 13.11) = 34,78%

Emmint	≥ " 1287	ntenes 46	Atomic Ratio	Empirical Formula
С	52.10	$\frac{52.10}{12} = 4.34$	$\frac{4.34}{2.17} = 2$	
Н	13,12	$\frac{13.12}{1.008} = 13.02$	$\frac{13.02}{2.17} = 6$	C₃H₅O
0	34.78	$\frac{34.78}{16} = 2.17$	$\frac{2.17}{2.17} = 1$	8

Thus the empirical formula for the organic compound is C. H.O

Example 5: The combustion analysis of an organic compound should be contain 65.445 carbon, 5.50% hydrogen and 29.06% of oxygen. What is the empirical formula of the compound? If the moiscular mass of this compound is 110.15. Calculate the moiscular formula of the compound is 110.15. formula of the compound.

(D.G. Khan Board, 2016: Foliolobed Boord, 2011: Gujranavala Board, 2005, 2007: Multan Board, 2016. Rayolpind Board, 2016, 2011)

Chambres Part-I

Solution:

ELIMINE		No. of grain	Atomic Ratio	Empirical Formula
С	65,44	$\frac{65.44}{12} = 5.45$	$\frac{5.45}{1.82} = 3$	
Н	5.50	5.50 1.008 = 5.45	$\frac{5.45}{1.82} = 3$	C _s H _s O
0 .	29.06	$\frac{29.06}{16} = 1.82$	$\frac{1.82}{1.82} = 1$	

Thus the empirical formula is C₃H₃O.

To determine the molecular formula, first calculate the empirical formula mass.

Empirical formula mass of $C_4H_5O = 12 \times 3 + 1.008 \times 3 + 16 \times 1 = 55.05 \text{ g mol}^{-1}$

Molecular (or moler) mass of the compound $= 110.15 \, \mathrm{g \, mol^{-1}}$

molecular mass of compound = $\frac{110.15}{55.05} = 2$

Molecular formula = n (empirical formula) Thus

$$= 2 \left[C_1 H_3 O \right]$$

$$= C_4 H_6 O_2$$

MOLE Exercise Q9: Define the following terms and give three example of each. (iv) Grom Ion (ii) Gram formula (ii) Gram atom (iii) Gram molecular mass

The atomic mass, molecular mass, formula mass or tonic mass of the substance.

Number of moles = Given Mess (in grams)

Examples:

Element

1 male of carbon = 12,000 g

Molecular compound:

1 mole of water = 18 g 1 mole of NaCt = 58.5 g

lente compound:

lonic species:

1 male ton of OH = 17 g

For each type of substance mole can be defined as

in grand is called one gra واله ويه الم محمد وا ole of that elame

Number of gram atoms or moles of an element — Mass of element in grams Asomic mass of an element

1 gram atom of hydrogen = 1.008 g

1 gram alom of carbon = 12,000 g

1 gram stom of Magnetium= 24 g

1 gram atom of Uranium = 238 g

Thus one gram atoms of different elements have different masses.

50, I atom of Mg is two times heavier than one atom of carbon

tream Make also Men Medecular compounds

The molecular mass of a substance expressed in grants is called one gran molecular one gran mole or steeply a mule of that substance.

Number of gram molecules or molecular substance in grams moles of a molecular substance in Mosecular mass of a molecular substance

1 gram molecule of water = 18 g I gram molecule of H_BSO₄ = 98 g

I gram molecule of sucrose= 342 g

Thus one upon molecule of different molecular substances have different masses.

Count Larged - if or foots comprounds

The formula mass of on tunta compound expressed in grame is called one god formula or one gram mais or simply a male of that substance.

Number of gram formulas or moles of an land substance in grams formula mass of an land substance in grams.

modes:

1 gram formula of NaCl = 58.5 g

1 green formula of AgNO₄ = 170 g

2 gram formula of Na₄CO₅ = 105 g

Charles Said

21

NOTE: funte compounds de not have molecular a

Since tonic compounds do not exist in malecular form, therefore, term matecular mass is not used for them. In lonk campaunds the sum of atomic masses of individual ions gives the formula mass

famo familiar basic species)

The tonic more of an tonic species expressed in grame to called one gram ion or one gram note or simply a mole of that for

Formula:

Number of gram lons ar males of an lonic species in grams Formula mass of an lonic species

Examples:

1 green ion of $OH^+ = 17 g$

1 gram ion of CO_1^{-1} = 60 g

1 gram tun of $SO_4^{2a} = 96 g$

Example 6:

.

Colculate the gram stains (moles) in

(a) 0.1 g of sodium

(b) 0,1 kg of silicon.

(a) 0.1 a of sodium

Given mass (in gracus) Number of gram atoms

> Mass of sodium - 0.1 g

= 23 g mpi 1 Moiar mass of sodium

Number of gram atoms of Na. = $\frac{0.1}{200}$ = 0.0043 moles = $\frac{4.3 \times 10^{-3} \text{ moles}}{1000}$ 23

(b) 0,1 kg of Silteon.

Mass of silicon $= 0.1 \text{ kg} = 0.1 \times 1000 \text{g} = 100 \text{g}$

Moler mass of silicon = 28.086 g mai-1

Number of green atoms = Given mass (in grams)

Molar ru

100 Number of gram atoms of St = $\frac{100}{28.086}$ = $\frac{3.56 \text{ evoles}}{}$

Culture Characterist Park-I

22

Sante Concesso

Emmedia 7

Calculate the mass of 10° moles of MgSO,

Solution.

Molar mass of MgSO_a

Moles of MgSO₄ = 10⁻⁸ moles

Number of moles of tonic substance = Given mass (in grams)

Motor mass

Given mass (In grams)

10-4 Thus

Mass = 120 x 10⁻⁹ = 0.12 g

Exercise Q9

Define the following terms and give three example of each

(vi) Avagadra s numbe

Acagadra's Na

It is the number of particles (atoms, Ions, malecules, formula units) present in one

Its value is 6.02×10^{13}

It is denoted by N_A

12 g of C = 6.02 x 10 m atoms of C = 1 mole of C

 $1008 \, g \, of \, H = 6.02 \times 10^{10} \, atoms \, of \, H$ = 1 mole of H

18 g of $H_4O = 6.02 \times 10^{28}$ molecutes of $H_4O = 1$ mole of H_4O

 $58.5 \text{ g of NaCl} = 6.02 \times 10^{10}$ formula units of NaCl=1 mole of NaCl

One male of different substances has different masses but some number of particles.

It is because <u>individual particles of different substances have different masses</u>, therefore, equal number of moles of different substances will also have different masses but same number of particles

- An atom of sodium is 23 times heavier than one atom of hydrogen. Thus in order to have same number of atoms, Na must be taken 23 times greater in mass than hydrogen.
- Similarly. Mg atom is twice heavier than C. Thus, 10 g of Mg and 5 g of C have same number of atoms.

Permulas to calculate the number of particles

Number of atoms, ions or molecules can be calculated by using following formulas.

lowic mest

Number of stores of an element α mass of the element α M_{α}

pricersic mass

Number of molecules of a compound = mass of the compound × Na Number of lone of sectionic specie - mass of like ton x Na

College Chemistry: Part-1

23

Boute Concepts

Simple Quick Calculation I xamples to Cuderstand Mole and Avoyadro's Number

- . One time water (H_zO) = 18 g \pm 6.02 \times 10⁴³ molecules Thus, one mole of water contains 2×6 02×10⁴⁵ atoms of hydrogen and 6.02×10⁴⁵ atoms of oxygen
- One mole of $H_9SO_4 = 98g = 6.02 \times 10^{25}$ molecules. Thus one male of HaSO4 contains 2×6 a2×10*3 atoms of hydrogen, 6.02×10*3 atoms of suphur and 4×6.02×10™ atoms of oxygen.

Example a.

Some substances ionize in water. Their number of positive and negative ions can be catculated.

Let 9.8 g of $H_1 SO_4$ has been dissolved in water

Number of moles of $H_aSO_4 = 9.8 \text{ / } 98\text{g} \pm 0.1 \text{ moles of } H_aSO_4$

H₄SO₄ ionizes in water completely

 $H_aSO_a \Rightarrow 2H^a + SO_a^{ab}$

Since, 1 mole of H₂SO₄ gives 2 moles of H² ions, 1 mole of SO₄ 2 ions, 2 moles of positive charges (two H^* wors, since each H^* wor has one positive charge) and 2 moles of negative charges (one SO_4^* rors, since each SO_4^* has two negative charges)

0.1 mole of H_aSO_a gives 0.2 males of H^+ ions, 0.1 mole of SO_a^+ ions, 0.2 moles of positive charges and 0.2 moles of negative charges.

Thus, Number of positive rous (H+ rous)

= 0.2×6.02×10*7 ions

Number of negative (ons (SO₄ * ions)

= 0.1×6.02×1043 tons

So, Total positive charges = 0.2×6.02×10¹³

Total negative charges = 0.2×6.02×10*3

Total mass of H+ .ons = (0.2×1.008)g

Total mass of SO_4 ions = $.0.1 \times 96$)g

Example 8:

How many molecules of router are there in 10.0 g of ice? Also calculate the number of atoms of hydrogen and oxygen separately, the total number of atoms and the covalent hands present in the sample.

(Gujronvels board, 2013)

```
College Chambers Parel
                                                                                                                                                                     2%
                                                                                                                                                                                                                Stanto Com-
                                                                                                                   (b) Number of socilies and operator loss in case of complete disconlation in senter
                           - 10 a
a. Meas of her (water)
                                                                                                                    ... H<sub>e</sub>PO<sub>4</sub> tonies in water as follows
   Moles mass all water = 18 g / mol

    Number of moles was of water = Given Mass (in green) = NA

                                                                                                                                          H<sub>1</sub>PO<sub>4</sub> --- 3H* + PO<sub>4</sub>*
                                                                                                                    - According to the balanced equation
                                                                                                                             1 molecule of H<sub>4</sub>PO<sub>4</sub> produces H' tons = 3
    Physicism of enchanges of unitary =\frac{10}{18} vis DSF a 10^{64} on 8.84 a 10^{68} production
                                                                                                                             6.14 \times 10^{10} molecules produce 14^{\circ} lore = 3 \times 6.14 \times 10^{19}
                                                                                                                                                                              =[1.042 × 10<sup>55</sup> lone
             arrota H antalago misso in alconduct
                                                             - 2
    3.34 \times 10^{16} \ \mathrm{molecules} of water contain 14 atoms . = 3 \times 3.34 \times 10^{16}
                                                                                                                     According to the balanced equation
                                                                                                                             ) implicable of H_aPO_a produces PO_a^{\ B} fone = 1
                                                              = 0.08 ×10<sup>64</sup> granu |
                                                                                                        )
                                                                                                                             6.14 \times 10^{16} molecules produce PO_a^{\, 6} sons = 1 \times 6.14 \times 10^{16}
              Annale Clarifolder retew to elevation 1
                                                                                                                                                                               -[6.14 \times 10^{27}]eng
     3.86 \times 10^{10} molecules of water contain O atoms. \approx 3.34 \times 10^{40} atoms
                  Total number of etoms of H and O _{\odot} = 6.68 \times 10^{14} \pm 3.84 \times 10^{18}
                                                                                                                     (a) Massas of Individual lone
                                                                                                                                       Number of limit = lonic mean of the ton
                                                                                                                                                               Mass of the ion
                                                               *140.08 × 1045 atomit
  If englands of water contain number of equalent bonds -- if
                                                                                                                                     Number of 14" lone - Mass of the 44" kins + MA
              3.34 x 10^{49} molecules of water contain _{\odot}=3\times3.34\times10^{18}
                                                                 Shoot Inslavue 40 to 98,0 p
                                                                                                                                            1 842 × 10<sup>10</sup> = Mess of the[-] some = 6 02 × 10<sup>23</sup>
                                                                                                                                                                     1.008
                                                                                                                                     | Mean of the H, town = \frac{2.08 \times 10_M}{1.948 \times 10_M} \times 1.008
  Kanstada U
  19.9 g of H_0 \!\!\!\! / \!\!\! G_0 have been disculsed in scenes of water in discontate () completely like two laborators
                                                                                                                                     Mean of the H1 tons = (0.894 at
   in Number of nuderates in 100 g of H_i P Q_i
   the blumber of position and sugation ions in case of samplists discussifing in center (1984)
                                                                                                                                                                                                lonic Moss of PIN lon - M. g.
                                                                                   (Ppleateber Double, 1944);
       to the number of $1" tone to $18 g of 14,1915,2
                                                                                                                                   Number of p \mathbb{Q}_q^{\frac{1}{p}} term = \frac{M_{\text{BM}} \circ l}{l \text{onto mass of } p \mathbb{Q}_q^{\frac{1}{p}}} torm
                                                                                                                      ,→ and
   (a) Massas of techniquel ture.
   Window of profitte and tragation charge dispersed its the solution.
                                                                                                                                                 5.14 × 10 Pt ~ Mass of the PCI4 1 John ~ 6.02 × 10 Pt
   Derbette
   ful Number of molecules in $8.0 g of $1,000,
                                                                                                                                                                            95
                                                                                                                                   Make of the PGq^1 -wire = \frac{6}{6}\frac{10.4}{10}\frac{10^{10}}{10} = 90
                         Mass of H_0 H G_0 = 1 G_0
                 Moles mass of H_0PO_0=3\in33+84=90(q\ell) and
                Number of melonates a Cityon mass (in grants) a NA
                                                                                                                                                                 -(9.069 g
                                                                                                                        (d) Number of position and resouther charges discurred in the authorism
                                                  Moles Hees

    I molecule of H<sub>0</sub>PO<sub>0</sub> produces positive changes = 3.

    Phimber of molecules of N_{e}PO_{ij} \approx \frac{10}{64} \times 6.08 \times 10^{10}
                                                                                                                                  9.14\times10^{10} molecules of H_0 PO_6 will give _{\odot}\approx3\times0.14\times10^{10}
                                                                                                                                                                                  - 1 848 = 10<sup>20</sup> | yaraktive changes
                                                          = 0.014 = 10<sup>55</sup> molecules

    Since Numbers of pusitive and negative charges are always equal.

                                                          =($.16 > 10<sup>10</sup> melecules
```

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Book Consequ

So the number of negative charges produced in the solution = 1.842 × 10²⁵

Ġ

Simuland Temperature and Pressure

Get volumes are usually compared at 0°C (273K) and 1 atm. These conditions are called as standard temperature and pressure (STP)

Exercise Q9

Define the following terms and give three example of each.

fu)Mator Volume

Malar Volume

The volume occupied by one main of an ideal gas at standard temperature and pressure (S.7 P.) is called main values and is equal to 22.414 dm³

The replan volume for real gases used for calculations is also 22,414 dm²

One mole of any gas contains constant number of molecules. This number is called Avoyadrus number is value is 6.02×10^{10}

Examples

I main at He = 6.02×10^{49} molecules of He = 2.016 g of H₂ = 22.414 dm² of H₂ at S.T.P. I make of CH₄ = 6.02×10^{38} molecules of CH₄ = 16 g of CH₄ = 22.4.4 dm² of CH₄ or S.T.P. I make of O₈ = 6.02×10^{38} molecules of O₈ = 32 g of O₉ = 22.414 dm² of O₉ at S.T.P.

22.414 dm² of each gas has different masses but some ramber of molecules.

It is because masses and sizes of molecules has no offect on volume of dases. In gases molecules are widely separated from one another and have large empty spaces. The distance between two molecules is approximately 300 times of the disameter of the

Excepte 10:

A well lose A well known ideal gov is enclosed in a container having column 500 cm² at S.T.P has make comes out to be 0.72g. What is the motor mass of this gas.

Volume of an ideal gas at STP = 500cm^b

Mass of ideal gas = 0.72 g

Moter mess of gas = ?

 $\star~22.414~dm^0$ or 22414 cm 0 of the ideal gas at S.T.P ≈ 1 mole

 500 cm^3 of the ideal gas at S T P = $\frac{1 \times 500}{200 \text{ keV}}$

22414

= 0 0223 males

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Basic Concepts

Number of motes = Given mass (in grams, Motor mass

Molar mass = $\frac{0.72}{0.0223}$ =[32 g / mol]

STOICHIOMETRY

Exarcles Q9:

Define the following terms and give three example of each full/Stoichiometry Exercise O.23, (a)

What is stoichlometery? Give its assumptions. Mention two important laws, which help to perform the stoichtometric colculations.

Definition: The branch of chemistry which deals with the study of relationship between the quantities of reactants and products as given by bolowerd chemical equation is called

Standiometric Assumptions

The stoichtometric calculations are based on following assumptions.

(I) Reactants are completely converted into products

(4) No side reaction occurs.

Lines to be obesed in Stoichnountre

For calculations, the law of conservation of mass and the law of definite proportions are obeyed

(i) Low of Conservation of Mass

It states

Massi can neither be created nor destroyed during a chemical reaction.

It can also be slated as

Total mass of reactants is aqual to the total wass of products.

Thus total number of atoms entering into a chemical reaction is equal to the total number of atoms in the products









2H, Q₂

24,0

Hants Concessor

(ii) Los of Dellaits Proportions

It states

ontains the same elements combined to the same A pure character comportation by weight.





Sect Appreciae Relationale pa

Moss-Moss Relationship

With the help of mass of given substance, mass of another substance can be calculated.

Moss-Mole Relationship

With the help of mass of one substance, moles of other substance can be calculated and vice verse.

Mass Volume Relationship

With the help of mass of one substance volume of the other substance can be calculated and vice versa.

Mote-Mote Relationship

With the help of moles of a given substance, moles of another substance can be calculated.

Advanceges and Livinstinus of Chronical Equations

Consider the reaction

$$C + O_2 \rightarrow CO_3$$

This equation tells that 1 mole of C, reacts with 1 mole of O_2 give one mole of CO_1 . Thus, this equation can be used to study quantitative relationship between reactants and products.

LinderHane

Chemical equations have certain limitations

- (I) These cannot beliabout the conditions of reactions
- (iii) These cannot tell about the rate of reactions.
- (III) These cannot tell about the time to complete the reaction.
- (M) These can also be written for such reactions, which are not possible

College Chemistry: Part-

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Stoichignearle amonnas The amount of react etalchicmetric amounts. ording to belanced chemical equation are called

Example.

$$C + O_k \rightarrow CO_k$$

This equation tells that 1 mole of C reacts with 1 mole of O_k . So, C to O_k mole ratio is 1.1 Any, amount which is according to this ratio will be stoichiometric amount: e.g. (12g C and 32g $O_{\rm B}$), (24g C and 64g $O_{\rm B}$) and (48g C and 128g $O_{\rm B}$, are all stoichiometric amounts.

Example II

Calculate the number of grams of K₂SO₄ and water produced when 14 g of ROH are reacted with assess of H₂SO₄. Also calculate the remoter of molecules of mater produced.

(D.G. Khan Board, 2011) Lehore Board, 2011 Multan Board, 2010: Sargodha Board, 3010)

Solution

The balanced chemical equation for the chemical reaction is

$$2KOH + H_2SO_4 \longrightarrow K_2SO_4 + 2H_1O$$

→ Given Mass of KOH = 14 g

Therefore

Molar mass of KOH = 39 + 16 + 1 = 56 g/mol

Motes of KOH =
$$\frac{14}{56}$$
 = 0.25 moles

According to balanced chemical equation

Compare the motes of KOH and KaSO.

KOH K₂50,

2 moles : 1 mole

0.25 moles . $\frac{1 \pm 0.25}{2} = 0.125$ moles

No, of moles of K_aSO_a produced = 0.125 moles.

Moler mass of
$$K_4SO_4 = 2 \times 39 + 32 + 4 \times 16$$

Thus Mass of KeSO₄ produced = No. of moles × Moles mass

For H₂O, compare KOH and H₂O according to the behanced equation.

Therefore 0.25 moles .
$$\frac{2 \times 0.25}{2} = 0.25$$
 moles

So the number of motes of water produced = 0.25 motes

Therefore Number of water molecules
$$= \frac{4.5}{18} \times 6.02 \times 10^{29}$$

$$= 1.505 \times 10^{49} \text{ malecules}$$

Example 12:

Mg metal reacts with HCl to give hydrogen gas. What is the minimum values of HCl (i) The recetant which consumes first in the chemical reaction is called limiting reactions.

Solution (27% by weight) required to produce 12.1 g of H_p. The density of HCl solution (9) The reactions which controls the amount of product formed during a chemical reaction is (Lohore Board, 2007, 2910) 1.14 gions

The balanced chemical equation for the chemical reaction is

→ Mass of H₆ produced = 12.1 g

Moler mass of H₂ = 2.016 g , moi

Moles of
$$H_0 = \frac{12.1}{2.016} = 6$$
 moles

Compare H_a and HCI according to balance equation.

He : HCI

1 mole . 2

Therefore 6 moles : $6 \times 2 = 12$ moles

Mass of HCl = Moles of HCl × Moles mass of HCl Hunes

4 12×35.5

438 g

→ Since solution of HCI is 27% by weight, therefore

31 27 g of HCl are present in HCl solution. = 100 g

100 - 438 438 g are present in HCl solution 27

= 16222g

→ Density of HCl solution = 1 14 g/cm² Mass of HCI solution = 1622 4 g

Since
$$d = \frac{m}{V}$$

or
$$V = \frac{m}{d} = \frac{1622 \text{ 4}}{1.14} = \frac{1423 \text{ cm}^3}{1}$$

1.14

a

LIMITING REACTANT

Exercise Q23 (b)

What is limiting reactant? How does it control the quantity of the product formed? Explain with three examples.

Definition

- called limiting reactant
- (b) The reactions, which gives least amount of product, is the limiting rengions. Explanation:
- When the reactants are not mixed in stoichiometric amounts, one of the reactants is consumed earlier. This is the limiting reactant. Other mactants are left unreacted.
- 1. When the limiting reactant is consumed, then reaction stops and further product cannot be produced. Thus, it controls the amount of product and gives the least amount of product

Example 1

$$2H_1 + O_2 \longrightarrow 2H_2O$$

When 2 moles of hydrogen (4g) reacts with 2 moles of oxygen (64g), then only 2 moles (35g) of water are produced. It is because, 2 moles (4g) of hydrogen react with 1 mole. (32g) of oxugen. Since, less hydrogen is present than oxygen, so hydrogen is the limiting reactant Promple 2:

Burning of coal occurs in excess of oxygen. In this coal is the limiting reactant while oxygen is in excess

Example 3;

Rusting of fron occurs in excess of oxygen present in air. In this, from is the limiting reactant while oxygen is in excess.

esten of Deliberate use of Limiting reactant

mple ! To completely renounts the expensive received

The other reactants are taken to large encore than expensive reactants. It ensures that a of the expensive renders in completely used up in the chemical reaction, so it becomes to limiting reactions.

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Energie R: To speed up the recotion-

Rate of reaction is directly proportional to the amounts of reactants. So, a large quant, of one reaction may speed up the reaction, e.g. a large quantity of oxygen burns thus faster. Thus excess of oxygen is left behind at the end of reaction and burning things to paper wood six are the limiting reacisms

Occumulation of Leading Resetable

The emount of product formed by a chemical reaction is calculated from the limiting reactant; therefore, identification of limiting reactant is necessary.

Following procedure is used for this

- (i) Number of moles of each reactant is calculated from given masses.
- 80 Using balanced chemical equation, amount of product, expected to form, from as martant is calculated.
- (te) The reactant, which gives least amount of product, is the limiting reactant

NH, gue can be proposed by heating together two solids NH_aCl and $Ca(OH)_b$. If suitable equivalency 100 g of each walld to heated them

- nists the number of grams of NHs, produced.
- (b) Coloniate the excess amount of reagent left unreacted.

Francistrat Board, 2010: Currenucie Board, 2011 Multan Board, 2013: Sargoche Sourd, 2011.

The balanced chemical equation for the chemical reaction is:

$$2 N H_s C I + C a (OH)_R \qquad \longrightarrow \ C a C I_R + 2 N H_R + 2 H_R O$$

miner of greens of KH₄ produced.

Given Mass of NH_eCl = 100g

Moler mass of NH_eCl = $14+1 \times 4+35.5 = 53.5 \text{ g/mol}$

Number of moles of NH_gCt = $\frac{100}{53.5}$

Given Mass of Ca(OH)_e = 100 g

Molar mass of Ca(OH); = 40+2(16+1 =74 g

Number of moles of Ca(OH)_{II} = $\frac{100}{74}$ = 1.35 moles

→ Compare NH₆Ci and NH₈ according to balanced chemical equation.

NH₄C3 : NH₄

2 moles : 2 males

 $\frac{2}{2} \times 1.87 = 1.87$ moles 1.87 motes Therefore

Compare Ca(OH)₄ and NH₂ according to balanced chemical equation.

Ca(OH)_a : NH_a

1 male : 2 moles

Therefore 1.35 moles : 2 ×1.35 = 2.70 moles

→ Since NH₄CI produces least amount of NH₂, hence NH₄CI is the limiting reactant. Thus no. of males of NH_a produced=1.87 moles

Molar mass of NH₅ = $14+1\times3=17$ g/mol

Hence Amount of NH_a produced = No. of moles × Molar Mass of NH_a

(ii) Amount of respent left unresoted.

→ Coropare NH₄Ct and Ca(OH)₈ according to balanced chemical equation

NH₄Cl : Ca(OH)_z

2 moles : 1 moles

1.87 moles : $\frac{1 \times 1.87}{2} = 0.935$ moles

Moles of Ca(OH)_s taken ≈ 1.35 mokes

= 1.35 - 0.935 = 0.415 moles Therefore unreacted moles

Mislar Mass of Ca(OH)_e # 74 g/mol

Thus Mass of Ca(OH)_e left = 0.415 x 74 = 80.71 g

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Exercise Q9:

Define the following terms and give three example of each. (viii)Percentage yield

Exercise Q.24 (a)

Define yield. How do we calculate the yield of a chemical reaction?

YILLD

The amount of product formed during a chemical reaction is called Yield.

Theoretical Vield

The amount of product calculated from beloaced chamical equation is called theoretical yield

It is the maximum yield of product that can be produced by a given amount of reaction a according to balanced chemical equation.

Example:

 $H_a \& O_a$ react together as

$$2H_s + O_s \longrightarrow 2H_sO$$

According to this equation 2 moles of H₂ & 1 mole of O₂ should form 2 moles of H₂0. Therefore, 2 moles of H2O is the theoretical yield

de tood Vector

The amount of product extently obtained during a chamical reaction is called Actal otald.

Percentage York for Lifticiancy of a Reaction)

The % yield or efficiency of the reaction is given by the formula

% Yield =

Exercise Q24 (b):

What are the factors, which are mostly responsible for the law yield of products in chemical reaction?

Actual yield of a chemical reaction is always test than its theoretical yield.

The reasons are

- (f) Inexperience worker wastes a significant amount of product.
- (iii) By product may be formed due to side reactions.
- (int) The reversible reactions are never completed.
- (Net Product is lost during their separation and purification processes, e.g. processes Stration, distillation, asparation by separating funnel, washing, drying and crystallization reduces the amount of product.

	Difference between Actual Yield and Theoretical Yield							
	Artual Yield	1	ti-rite-riski					
1	It is the amount of a product actually obtained during a chemical reaction.	1	It is the amount of a product calculated from a balanced chemical equation.					
2	It is always less than the theoretical yield	2	It is always greater than actual yield.					
3	Reversible nature of the reaction and side reactions decreases the actual yield.	3	White calculating theoretical yield, it is assumed that (i. No side reaction occurs (iii) Reactants are completely converted into products. Therefore, it is not decreased.					
4	Loss of product during separation, purification and by inexperience worker decreases the yield.	4	No mechanical losses occur, since it is calculated theoretically.					
5	if is an experimental form.	5	It is a theoretical term					

Example 14:

When time stone (CaCD₂) is rounted, guiokline (CaO) is produced occording to the following equation. The actual yield of CaO to 2.5 kg, when 4.5 kg of time stone to rounted. What is the percentage yield of this reaction?

(Fulsalabad Board, 2013: Multan Board, 2010: Rauselpinet Board, 2013)

Solution

The balanced chamical equation for the chemical reaction is

$$CaCO_3 \longrightarrow CaO + CO_4$$

- $= 4.5 \, \text{kg} = 4500 \, \text{g}$ Mass of lime stone
 - Moter mass of CaCO_a = 100 g/mol
 - = 4500 / 100 = 45 moles Moles of CaCO₂ *
- → Mass of quick time produced = 2.5 kg = 2500 g (actual yields)
- → According to the belanced chemical equation

Compare the moles of CaCOs and CaO

CaCO, . CaO

1 mole : 1 mole

Therefore 45 males : 1 x 45 = 45 males

Moler mess of CeO = 40+16=56 g/mol Think Mass of CaO produced = 46 x 56 × 2520 g Hence Theoretical yield of CaO = 2520 g Actual yield of CaO = 2500 g % yield is given by Actual yield % yield = Theoretical yelld × 100 $\frac{2500}{2520} \times 100$ =99.2%

37

OPTICATION AND SHORT ASSECTE OF ESTAGES (USED INC.)

- Select the most suitable master from the given cross to each question
- (1) leotopes differ le
 - (a) properties which depend upon mean
 - (b) arrangement of electrons in orbital
 - (c) chemical properties
- (d) the extent to which they may be affected in electromagnetic field

dead Board, 2009. 2011. D.G. Khan Board, 2009. Fatesballed Board, 2011. D.G. Khan Board, 2012)

- (N) Which of the following etalement is not true? (Labora board, 2013)
 - (a) isotopes with even atomic masses are comparatively abundant (b) isotopes with odd atomic masses are comparatively abundant.

 - (c) isotopes with even atomic masses and even atomic numbers are comparatively abundant.
- (d) isotopes with even atomic masses and odd atomic numbers are comparatively ebundanı
- (iii) Many elements have fractional attente mesors, this to because
 - (a) the mass of the atom is itself-ractional
 - (b) atomic masses are average masses of isobara
 - (c) atomic masses are average masses of botopes
 - (d) atomic masses are everage masses of isotopes proportional to their relative abundance

(bc) The means of one mote of electrone is

(a) 1.008rng (b) 0.55rng (c) 0.189rng (d) 1.673rng
(Febolahod Board, 2007, 2009, Mallon Board, 2012, 2012, Surgadha Board, 2012, 2014, Casternatolo Board, 2010, 2012, 2015, Labore Board, 2014)

(v) 27 g of Al will react completely with how much mass of O₂ to pre-duce Al₂O₅
(a) 8g of oxygen (b) 16g of oxygen (c) 32g of oxygen (d) 24g of oxygen
(44) 24g of oxygen (b) 16g of oxygen (d) 24g of oxyge

(vi) The number of moles of CO₄ which contain 8 g of sayyon (a) 0.25 (b) 0.50 (c) 1 (d) 1.5

(Roundplant Bourd, 2009: Gujermeele bourd, 2009, 2018: 2014: Multen Bourd, 2013)

(vii) The largest number of molecules is present in

(a) 3.5g of H₂O (b) 4.5g of C₄H₂OH (c) 2.8g of CO

(A) 3.5g of H₂O (c) Non-Board, 2009 Suspecte Sound, 2011, 2012; Resolved Sound, 2011; Males Sound, 2011; Lateur Board, 2014)

(viii) One mole of SO_2 contains (a) 6.02×10^{33} alone of oxyger

(b) 1.81×10^{39} molecules of $5O_2$

(c) 6.02 x 10⁴⁴ etems of sulphur

(d) 4g atom of SO₂

Pipur Board, 2009: Sergodhe Board, 2009: Cultoruelo Board, 2013: Multer Board, 2013;

(b) The solutions occupied by 1.4g of N_g at STP is
(a) 2.24dm⁸ (b) 22.4dm³ (c) 1.12dm⁸ (d) 112cm⁸

(Garranuccio hossel, 2008, 2014, Sargodhe Board, 2009, 2011, Lahans Soard, 2010)

3.3

- A function remotest is the one which

 (a) a taken in issuer quantity in grams as compared to other reactants

 (b) is taken in lesser quantity in volume as compared to the other

 (c) carries the maximum amount of the product which is required

 (d, gives the minimum amount of the product under consideration

made board, 2009: Multon Board, 2010: Behavelper Board, 2010)

ANSWERS DEMOCRATE	choice on AlloNa
Inchopes have some phonic number but different	in New skill holizon with odd oxenic manyly are comparatively and alumdant.
eriocolis, massisti.	technique with even accente number and even most are note abundent Out of 280 natural schools 164 have sign mass number and quiet anomic number.
The storile reviews of elements are calculated from their partiest of storopys and their patient shandards. Since the calculated resistant are made interested the calculated resistant are sounds. Theses, therefore, many elements have freeweeps according to the calculated resistants.	Fig. 3.7 in the residue $= 0.02 \times 10^{13}$ slectrons $= 0.02 \times 10^{13} \log 10^{13} \log 3$, electron than mining $= 9.1 \times 10^{13} \log 3 \times 10^{13} \log $
L. Maximilia	32 g. of O are present to CO ₂ = a mole
Make of $Al = \frac{27}{27} = 1$ mole According to behanced eq.	B_{\parallel} at O are present in $CO_3 = \frac{1 \times B}{32} = 0.25$ mole
ACCOUNTS OF STATE OF ACCOUNTS	
And America	
No of protection of $H_0O = \frac{3.5}{18} \times 6.02 \times 10^{10}$ = 0.2 \times N _c	One most of SO ₂ contents: Doe mode of Sulphus storms, and Two moins of oxygen storms Thus, No. of sulphus abone 1 k 6.02 k 10 ⁹⁶ and
No. of molecules of $C_aH_aGH = \frac{4.8}{46} \times 4.02 \times 10^{64}$ =0 $_2 \times H_a$	No of Oxygen Atoms = 2x 5.02 x 10 th
No of molecular of QO = $\frac{3.4}{26} = 6.02 \times 10^{49}$ =0.1 × N _a	
No. of molecular of N ₀ O ₀ = $\frac{6.4}{106} \cdot 6 \times 10^{20}$ = 0.05 N ₁	
the later of	Oct Yes all
Holes of $N_0 = \frac{1.4}{28} = 0.05$ main	The reaction which gives minimum errorest product is called limiting reaction.
. maior of N _e at S.T.P. conspine you =22.4144gg ² 0.05moin N _e at S.T.P. conspice you =22.414d0,05 =1.12 dm ²	_ <u></u>

2.9	(i)	In the blanks The unit of relative atomic mass is expressed in
	(0)	The exact masses of isotopes can be determined by
	(60)	The phenomenon of isotopy was first discovered by
	(iv)	Empirical formule can be determined by combustion analysis for those compounds which have and in them.
	(v)	A similing reagent is that which controls the quantities of
	(v)	$1\ \mathrm{mole\ of\ glucose\ has}$ atoms of carbon, of oxygen and of hydrogen
	(VII)	4 g of CH ₄ at 0°C and 1 atm pressure has molecules of CH ₄
	(viii)	Strichtometric calculations can be performed only whenis obeyed.
	viil) lau	infact shady (eli 6 N., 12 N., 6 N., 1905) (viii. 1.505 x 10 ^{ca}) of conversation of mass deate may be Near has three isotopes and the fourth one with atomic mass 20 18 arms.
	(11)	Empiritized formula gives the information about the total number of atom- present in the molecule.
	(10)	During combustion analysis Mg(ClO ₄) ₈ is employed to absorb water vapours.
	(fu)	Molecular formula is the integral multiple of empirical formula and the integra multiple can never be unity
	(v)	The number of atoms in 1.79g of gold and 0.023g of sodium are equal.
	(vi)	The number of electrons in the molecules of CO and $N_{\rm B}$ are 14 each animg of each gas will have same number of electrons.
	(vii:	Avogadro's hypothesis is applicable to all types of gases i.a. ideal and nonideal
	(viii)	Actual yield of a chemical reaction may be greater than the theoretical yield

(68) True (cdif)false

(i) Positive ions are produced by passing high energy electron beam, or particles or X-rays through a gas.

Cone are those species which carry either positive or negative charge.

Q4. What are jone? under what conditions are they produced?

These may be produced

40

No + No' + e' e.a.

(ii) Negative love are produced by the addition of an electron to a neutral species.

CI+e + CI e.Q.

(MI) By ionization of an jonic compound in water.

e.g. NaCl = Na' + Cl

Q5.(a) What one the leatopes? How do you deduce the fractional atomic mass elements from the relative isotopic chundances? Gipe and examples in the support of

Salved on Page

(b) How does a more spectrograph show the relative abundance of isotoper of an

Solved on Page 12

(c) What is the fustification of two strong peaks in the mass spectrum of bromine, while for loding only one peak at 127 a.m.n. is indicated?

Bromine has two neturally occurring isotopes with aimost equal relative abundance. therefore, its mass spectrum shows two strong peaks.

While indine has only one naturally occurring isotope, therefore, its spectrum shows only one peak.

Q9. Duffine the following terms and give three example of each.

(i) Gram atom: Solved on page 20

(8) Gram molecular mass: Solved on page 20

(ii) Gram formula: Solved on page 20 (v) Molar Volume: Solved on page 26 (iv) Gram ion: Solved on page 21

(vii)Stoichlometry: Solved on page 27

(vi) Avogadro's number: Solved on page 22

[vili]Percentage yield: Solved on page 34

Q9. Justify the following statement,

(a) 23 gram of socition and 288 grams of translate hope equal number of atoms in them. (Faladabad Bord, 2007; Labors Board, 2006; Mukim Board, 2010)

1 mole of Na = 23 g

 $1 \mod old U = 238 g$

Since 1 mole of each element contains Avogadro's number of atoms and there is 1 mole of each of Na and U. Hence 23 g of Na and 238 g of U contains equal number of atords. La

(b) Mg atom is turice hemier than that of carbon.

(Gujimanaka Board, 2011, 2014). (Labors Board, 2007, 2011), (Rassalphel Board, 2008, 2010) One atom of cashon contains 6 protons and 6 neutrons in its nucleus and its atomic mass on atomic mass unit scale is 12 arms.

POLYTHAN CY / THE CONTRACT OF THE CONTRACT OF

While one atom of Mg contains 12 protons and 12 neutrons in its nucleus and its attents coass or atomic mass unit scale is 24 amu. Thus, 24 amu = 2. Hence, one atom of Mg at the housing than that of one of Mg at twice heavier than that of one atom of carbon

(c) 180g of glucose and 342g of sucross have some number of moleculus but different of atoms. (Gutumunia Board, 2012: Lehore Board, 2006: Sargodia Board, 2012)

180 g of glucose ≠1 male

342 g of sucrose = 1 mole

1 mole of each compound contains Avogadro's number of molecules. Hence, 180 g of glucase (1 mole) and 342 g (1 mole) of sucrose contain equal number of molecules. i.e. 6.02×10²³

Since one molecule of glucose (C_eH₁₂O_e) contains 24 atoms. Whereas one molecule of sucrose $(C_{12}H_{EG}O_{12})$ contains 45 atoms, therefore, equal number of molecules of glacose and sucrose will have different number of atoms.

(d) 4.9g of H₂SO₄ when completely located in water have equal number of position and negative charges but the number of positively charge loss are taken the number of negatively charge tone. (Lahore Board, 2012)

H₂SO₄ ionizes in solution as

H₂SO₄ ≠ 2H* + SO₄**

This balanced equation shows that

1 malecute of H₂SO₄ produces

Number of positively charged ions (H* ions) = 2

Number of negatively charged ions (SO₄*) = 1

Number of positive charges = 2 (due to two H* ions)

Number of negative charges = 2 (due to two negative charges on 50.4)

Hence, whatever be the amount of H₂SO, it will always produce agual number of Positive and negative charges but number of positively charged ions will be twice the

number of negatively charged ions.

(e) One mg of $K_0Cr_2O_1$ has thrice the number of low than the number of molecules when ionised in conter. (Labore Board, \$010: \$012, 2013)

K_pCt_pO_p ionizes as

 $K_0Cr_2O_7$ = $2K^4 + Cr_2O_7^{2-}$ This equation shows that 1 formula unit of $K_0Cr_2O_7$ produces two K^4 ions and one Cr₂O₇² ion in solution. Thus a total of three ions are produced by the ionization of 1 formula unit of KaCraO2.

Hence whatever be the assount of $K_{\theta}Cr_{\theta}O_{\phi}$, number of ions in its solution will always be finite than the number of its formula units (molecules)

(f) Two presss of H₂, 16g of CH₄ and 44g of CO₂ accepts separately the volume of 22.414 del alchem**y** distances of molecular of these games are very different from (Multon Bornd, 2007) party other

2 grams of $H_R = 1$ mole = 6.023×10^{43} molecules

16 grams of CPI₄ = 1 mole = 6.023 × 10²⁴ molecules 44 games of CO₅ = 1 mole = 6.023 × 10²⁴ molecules

Since 2 g of H_B 16 g of CH_b and 44 g of CO₂ contains equal number of molecules and according to Avegadina's law, equal molecules of all gases at S.T.P. occupy same votume re

22 414 der in gases distance between two molecules is approximately 300 times its molecular size Thus, volume occupied by gas molecules does not depend upon the size or mass of molecules and it only depends upon the number of molecules.

Hence equal moteraists of $H_{\rm b}$, $CH_{\rm c}$ and $CO_{\rm c}$ at STP will occupy same volume \approx 22 414 dm³

Q23 (a) What is stoichismetery? Give its assumptions. Mention two important lass, thich help to perform the stoichiometric calculations

(b) What is limiting recetant? Hose does it control the quantity of the product formed? Explain with three exemples.

Solved on Page 31

Q24 (a) Define yield. How do we calculate the yield of a chemical reaction?

Solved on Page 34

(b) What are the factors, which are mostly responsible for the low yield of products in chemical reaction?

Solved on Page 35

430

Q25 Explain the following with reason.

(I) Les of conservation of most has to be considered during stoichiometric calculations sold (Faisalabad Board, 2007: Lahure Board, 2013)

Stoichiometric calculations are done with balanced chemical equations in which it is antumed that realist is neither created nor destroyed during a chemical change. Hence law of conservation of mass has to be considered during stoichiometric casculations, otherwise, with unbalanced equations stoichlometric calculations will not be possible.

4.5

(ii) Many charactel reactions taking places in our surrounding insche the limit

A limiting seactant is one which has limited quantity and consumes first in a chemical

There are many chemical reactions in our surroundings which involve limiting reactions

Petrol burns in sucess of oxygen present in sir

Rusting of son occurs in excess of oxygen present in air

Burning of roal occurs in excess of oxygen

in above cases, petrol, from and coal are limiting reactant while oxygen is in excess

(8) No individual new atom in the sample of the element free a more of 20.18 mms. (Gajanusia Board, 2013: Latera Board, 2014: Multan Board, 2007, 2012. Surgothe Board, 2013)

A sample of neon consists of $^{20}_{10}$ Ne, $^{21}_{10}$ Ne and $^{22}_{10}$ Ne in the percentages of 90 92% 0.26%, 8.82% respectively. The average relative mass of Ne is thus calculated as

average atomic mass =
$$\frac{(20 \times 90.92) + (21 \times 0.26) + (22 \times 8.82)}{100} = 20.18$$

Hence 20.18 amu is the average atomic mass of neon and no individual mean atom in the sample that has a mass of 20 18 arm.

(iv) One make of H₂SO₄ should completely react with two moles of NaOH. How done dro's number help to suplain it (Roselpind Board, \$606, 2009)

The chemical reaction of H₈SO₄ and NaOH is H₂SO₄ Indepute 2NoOH * 2 formule units \$ molecule ov 2 × 5 02 × 10th formula units 6 02 × 10th mole

NagSO₄ + H_zO

Hence 2 males of NaOH reacts with 1 mole of H₂SO₄

(a) One make of H₂O has two makes of band, these makes of stome, test makes of electrons and intensity-eight moles of the total fundamental particles present in it.

I molecule of H₂O contain bonds

6.02×10^{ss} molecules contain bonds = 2×6.02×10^{ss}

Thus 1 mole of H_aO contain bonds = 2 moles

I molecule of H_sO contain atoms

6.02 x 10^{2a} molecules contain atoms = 3 x 6.02 x 10^{aa}

Thus 1 mole of H_sO contain atoms = 3 moles

Dectrons

I molecule of H_EO contains two H atoms and one O alom.

```
To find the Tage of element in a composition of the planning o
  6 02 x 10<sup>nd</sup> molecules contain electrons = 10 x 6.02 x 10<sup>nd</sup> molecules contain electrons = 10 moles
    of Fundamental Furticles

1 majorate of HyO contemps two H atoms and one O atom.
                                                                                                                                                                                                                                                     Hed the Frenchical Formula
    Smt#
              1 O atoms contain electrons—8, Protons—8 and Neutrons—8
              1 () atoms contains total fundamental particles = 8+8+8=24
       Serios
               $ 14 alone contain electrone=1, Protons=1 and Pleutrone=0
                                                                                                                                                                                                                                                                                                             A 166 A
                                                                                                                                                                                                                                                                     Thought Makes regard on some Vis
                                                                                                                                                                                                                                                                      Region Rose & glass of 25 g
                1 H streng contine total fundamental particles = 1+1+0=2
                2 H annes systalis total fundamental preticles = 4
                                                                                                                                                                                                                                                                                                                                                                                           14
                                                                                                                                                                                                                                                         A Surger F. S. Later Sections
                 1 materials of H<sub>2</sub>O contain fundamental particles = 24+4=28
                4.62 × 10<sup>th</sup> nucleocides common hundersemial particles = 20 · 6.02 × 10<sup>th</sup>
Thus 1 mole of H<sub>2</sub>O contain bonds = 28 moles
                                                                                                                                                                                                                                                                   PEACL, Eux cames objectioned more passes seen on sounds
                                                                                                                                                                                                                                                 To flad projecular formula from empirical formula
tell H<sub>2</sub> and CO has come number of electrons, protons and nustrons 4./5 him board 2009 Feinhald Board, 2010, 2012 Geforeude Board, 2006, 2012, 2015. Stiller Board, 2012 Seguelle Board, 2011)
                                                                                                                                                                                                                                                                             Microscot mass - Globocular formula = n - empetod formula:
Respectors sortische Illand
The elements C, H and O are represented as $2C, $4N and $2O
                                                                                                                                                                                                                                                 To find number of ancies or intee or paylor mass of a compound
                                                                                                                                                                                                                                                                                              Miches Human
Thus
                           1 molecule of N<sub>2</sub> contains electrons = 7+7=14 (7 for each N atom) 1 molecule of CQ contains electrons = 6+8=14 (C=6 , O=8)
                                                                                                                                                                                                                                                                     blotes Many many by Modestales more on elempts, more on formula more on jurils, mass
                   similarly.
                                                                                                                                                                                                                                                  To find the number of paritains (stome, tom, melecules or formule units)
                           1 moderate of Pt<sub>a</sub> contains protons = 7 + 7 = 14. (7 for each N atom. 1 moderate of CO contains protons = 6 + 8 = 14. (C=6, O=8)
                                                                                                                                                                                                                                                    No oil granticine = Mose = 94, or No oil preticine = moles = 14, 
Motor Mose = 94,
                  and
                           I molecule of N<sub>3</sub> contains neutrons = 7 + 7 = 14 (7 for each N etom). I molecule of CO contains neutrons = 6 + 8 = 14 (C = 6 + O = 8).
                                                                                                                                                                                                                                                                     Mader More may by Mulecular mass or stands mass of familia mass or some mass
                                                                                                                                                                                                                                                  To find the Kegg yield or efficiency of the searches R_0 yield \sim \frac{R_0 t_{\rm bol} \cdot V_{\rm bol}}{Theorem, a Yield} x (00)
```

ē t

Sende

One O atoms contain electrons = 8 One H stome contain electrone = 1. Two H atores contain electrons = 2

not I molecule all H₂O contain electrons. = 2+6=10

College Chemistry Fort-I

Baric Conoceta

45

IMPORTANT FORME LAS To find the overage about more and elementation masses also upopes and lead read in about the Re-Megragetorhiches Passope - Sabundaroe - pussoi d'asocope - Sabi ndance .

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Calless Chemistry: Part-1
                               NUMERICAL PROBLEMS (Exercise)
     Silver has atomic number 47 and has 16 known isotopes but only top occu-
naturally i.e. <sup>100</sup>Ag and <sup>100</sup>Ag. Given the following mass spectrometric data.
Calculate the overage atomic mass of affiner.
                                                                                                                                  (b) Males of O stome in 9.00g of Mg(NO<sub>3</sub>),
                                                                                                                                                                                                                     (Sergodia Board, 2013)
                                                                                                                                                            Mass of Mg(NO<sub>4/2</sub> = 9g
                                                                                                                                                  Molar Mass of Mg(NO_a)_g = 24 + 14 \times 2 + 16 \times 6 = 148 \text{ g mol}^{-1}
                                            Percentage Abundance
    sotopes
                   Mass .a.m.u.l
                                                                                                                                            Number of moles of Mg(NO_9)_8 = \frac{G_1 \text{ver mass (in grams)}}{2}
                                                                                                                                                                                         Molar mass
9
                    106 90509
                                             51.84
        'Ag
    MAg
                                              48.16
                    108 90476
                                                                                                                                                                                      = 9 ± 0.0608 moles
Solution:
                                              (106 90509 × 51 84, + (108 90476 × 48.16)
        Average atomic mass =
                                                                                                                                               more of Mg(NO<sub>3</sub>)<sub>2</sub> contains O motes
                                                                           100
                                                                                                                                                                                                           = 6 males
                                                                                                                                             0.0508 moles of Mg(NO<sub>1/4</sub> contain O moles = 6 \times 0.0608 = 0.3648 males
                                         = [107.87 q.m.q.
                                                                                                                       Q.
                                                                                                                                   (c) Number of O alone in 10.037g of CuSO<sub>2.5</sub>H<sub>2</sub>O
Q7 Boron with atomic number 5 has two naturally accurring feetopes. Calculate the
percentage abundance of "B and "B from the following data;
                                                                                                                                                       Mass of CuSO_4 5H_2O = 10 037 g
                                                                                                                                             Molar Mass of CuSO<sub>4.5</sub>H<sub>E</sub>O = 63.5+32+1\times10+16\times9=249.5 g mol <sup>4</sup>
        Average Atomic mass of B
                                                    = 10.81 amu
                Isotopic mbss of <sup>10</sup>B
                                                    = 10.0129 \text{ amu}
                                                                                                                                            Number of moles of of OuSO<sub>4</sub>,5H<sub>2</sub>O = Given Mass
                leotopic mass of <sup>LL</sup>B
                                                    = 11.0093amu
                                                                                                                                                                                                     Motar Mass
 Solution
                                                                                                                                                                                                     10.037
                      % abundance of <sup>10</sup>B ⇒ x
                                                                                                                                                                                                     249.5
            Then % abundance of ^{13}B = 100 - x
                                                                                                                                                                                                 = 0.04 moles
            Thus
                                                                                                                                                  1 mole of CuSO<sub>4</sub>.5H<sub>2</sub>O contain oxygen = 9 moles 0.04 moles CuSO<sub>4</sub>.5H<sub>2</sub>O contain oxygen = 0.04 \times 9 \times 0.36 moles
          Average atomic mass = \frac{[10.0129 \times x] + [11.0093 \times (100 - x)]}{2.25} = 10.81 \text{ and}
                                                               100
                                                                                                                                                                                            = 6.02 × 10<sup>23</sup> atoms
                                                                                                                                                  1 more of oxygen atoms
                                      \alpha = 10.0129 x + 1100.93 - 11.0093 x = 1081
                                                                                                                                                  0.36 males of oxygen atoms = 0.36 x 6.02 x 10<sup>ns</sup> atoms
                                      or -0.9964 \, x = 1081 - 1100.93
                                                                                                                                                                                            = 2 167 × 10<sup>22</sup> atome
                                           -0.9964 \times = -19.93
                                                                                                                                   (d) Mass in kilograms of 2.6 × 10<sup>th</sup> molecules of SO<sub>T</sub>
(Folsolabed Board, 2010: Gujrams
                                                        x = \frac{19.93}{0.9964} = 20,002\%
                                                                                                                                                        Number of molecules of SO_2 = 2.6 \times 10^{29}

Molar Mass of SO_2 = 2.6 \times 10^{29}

Molar Mass of SO_2 = 32 + 16 \times 2 = 64 \text{ g mol}^{-1}
                                                                                                                                                                                                                      rrd, 2006: Labore Sourd, 2014)
             Hence % abundance of ^{10}B = x = 20.002\% and % abundance of ^{11}B = 100 - 20.002 = 79.998\%
                                                                                                                                              Number of molecules of SO<sub>2</sub> = Given mass (in grams) × N<sub>a</sub>
                                                                                                                                                                                                  Motar mass
   Q.10 Calculate each of the following quantities.
                                                                                                                                                                   2.6 \times 10^{10} = \frac{\text{Mass}}{6.4 \text{g}} \times 6.02 \times 10^{23}
   (a) Mass in grows of 2.24 moles of HMnO (Multan Board, 2012, 2012; Sargedha Board, 2010)
                                                                                                                                                                          Mass = \frac{2.6 \times 10^{20} \times 64}{6.02 \times 10^{23}} = 2.76 \times 10^{-9} \text{g}
                  Moles of KMnO<sub>4</sub> = 2.74 mol

Moler Mass of KMnO<sub>4</sub> = 39+55+16 \times 4 = 158 g mol<sup>-1</sup>
                                                                                                                                                                                        6.02×10<sup>23</sup>
                  1 mole of KM<sub>2</sub>O<sub>4</sub> = 158 g
2 74 oles of KM<sub>2</sub>O<sub>4</sub> = 2 74 × 158 = 432,92 g
                                                                                                                                                                   Mass n kg = 2.76 × 10<sup>-2</sup> = 2.76 × 10<sup>-6</sup> kg
                                                                                                                                                                                       1000
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Basic Column

46

College Chemistry: Part-I

College Chaptern: Partil (Surpodha Board, 2011) (e) Moles of CI atams in 0.822g C₂H₄Cl₂ = 0.822 g = $12 \times 2 + 1 \times 4 + 35.5 \times 2 = 99 \text{ g mol}^{-1}$ Mass of CaHaCla Motor Mass of CallaCla Number of males of $C_2H_4G_3$ # Given mass (in grams) Number of moles of $C_3H_4Cl_g=\frac{0.822}{\alpha\alpha}=8.303\times10^{-5}\,\mathrm{moles}$ 1 mole of $C_0H_4CI_c$ contains CI atoms =2 moles 8.303×10^{-8} moles contain CI atoms $=2\times 8.303\times 10^{-8}$ = 0.0166 moles (f) Moss in grams of 5.136 moiss of effect corbonate ð Moles of $Ag_kCO_a = 5.136 \text{ mol}$ Moles Mass of $Ag_kCO_a = 275.74 \text{ g mol}^{-1}$ Number of moles of Ag₂CO₃ = Given mass (In grams) $5.136 = \frac{Main}{275.74}$ Mass Mass = 5.136 × 275.74 = 1416.2 g (g) Mose in grome of 2.78 × 10th molecules of CrO_xCl_x $= 2.78 \times 10^{83}$ Number of molecules of CrO₂Cl₄ $= 52 + 16 \times 2 + 2 \times 35.5 = 155 \text{ g mol}^{-1}$ Molar Mass of CrO_gCl₂ Number of molecules of $CrO_{g}Cl_{g} = \frac{Given meas (ln grams)}{N_{A}} \times N_{A}$ Moler mess $= \frac{M_{635}}{155} \times 5.02 \times 10^{23}$ 2.78× 10th 6.02×10^{23} (b) Number of moles and formula units in 100g KCiO Mass of KCIOs = 100 g Molar Mass Of KCIO₈ = 39+35.5+16×3=122.5 g mol -1 _ Given Mass (in grams) Number of moles of KCIOs Molar mass

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<u>College Chamberry Period</u> $=\frac{100}{122.5}$ = 0.816 moles Given Mass (in grams) × Na Number of formula units Moler mass $= \frac{100}{122.5} \times 6.02 \times 10^{23}$ = 4.91 × 10²⁷ formule unite (i) Number of K* ions ClO₂* ions, Cl atoms, and O atoms in (h)

Formula unit of KClO₂ = 4.91 × 10⁸³ 1 formula unit of KClO₃ contains K* tons = 3 4.91 × 10^{ss} formula units contain K* ions = (4.91 × 10^{ss} ions 1 formula unit of KCiO₃ contains CiO₃* tona= 1 4.91 \times 10²⁸ formula units contain CiO₃* tona= 4.91 \times 10²⁸ formula units contain CiO₃* tona= 1 formula unit of SCIO₃ contains Cl atoms = 1 4.91 × 10⁴⁴ formula units contain CI atoms= 4.91 × 10⁴⁴ atoms 1 formula unit of KCIOs contains O atoms = $3 \times 4.91 \times 10^{23}$ formula units contain O atoms = $3 \times 4.91 \times 10^{24}$ = 1.478×10^{24} atoms 911 Asportume, the critical sweetner has a molecular formulae of $C_{iq}H_{iq}N_{i}O_{q}$ (e) What is the mass of one mole of aspartome $(C_{ij}H_{ij}N_{ij}O_{ij})$ Mass of one mole of aspartame = $(12 \times 14) + (1 \times 18) + (14 \times 2) + (16 \times 5)$ - 294 g a (b) Hose many moles are present in 52g of aspartance Number of moles = Given Mass (in grams) Molar mass = 52 294 = 0.177 moles

College Chemistry Part f 51 (b) Which element is represented by the symbol 14? Mass of MF₂ produced Moles of MF₂ produced = 0.600 moles 0.600 moles of MF₁ = 46.8 g 1 mole of MF₂ = $\frac{46.8}{0.600} = 78g$ Formula mass of MF $_{\rm II} = 78$ Alomit mass of M + (19 \times 2" = 78 Atomic mass of M = 78 - 38 = 40Since atomic mass of Ca is 40, therefore M is Ca. Q15. In each pair choose the larger of the indicated quantity, or state if the samples are equa (d) Individual particles: 0.4 males of august molecules or 0.4 atoms anygen stores.
(b) Mass: 0.4 male of ozone molecules or 0.4 males of axygen atoms (c) Mass: 0.6 mole of Call, or 0.6 moles of I, (d) Individual particles: 4.0g NaO, or 3.3g SO, (8) Total ions: 2.3 males of NaCIO, or 2.0 males of MgCI, (f) Molecules: 11.0g H₂O or 11.0 g H₂O₂ (g) Na* ton: 0.500 moles NaBr or 0.0145 kg NaCl (h, Mass: 6.02 × 10²⁵ stoms of ²⁵U or 6.02 × 10²⁵ aloms ²⁸⁸U Both have same number of particles Since both are equimpolar quantities, therefore they $L_0.04 \times 6.02 \times 10^{23} = 2.408 \times 10^{23}$ particles have equal number of perpotes. For O, hor D Moles of O₃= 0.4 moles Mo.es of O = 0.4 Moles mass of $O_0 = 48 \text{ g mol}^3$ Mass = moles × moles mass Atomic mass of $O = 16 \text{ g mol}^3$ Mass = moles × atomic mass

= 0.4 x 16 a 6.4 g ater mass than 0.4 moles of O

Number of moles = 0.6

Mass = moles × motor mass

Mass of I_a = 0.6 × 253.8

rdı

Маке и 50,

= 152 28 g

ter mass than 0.6 males of C.H.

Motor mass of SO₈ = 32+32 = 64 g most

Motor mass of $l_0 = 2 \times 126.9 = 253.8 \text{ g mpl}^4$

= 04 × 48 = 192 g Hence, 0.4 moles of O, has gre

 $= 0.6 \times 28$

Mates of $N_g O_q = 4 g$ Motor mass of $N_g O_q = 28 + 54 = 92 g mol$

= 16.8 g Henre, 9.6 males of 1, has gr

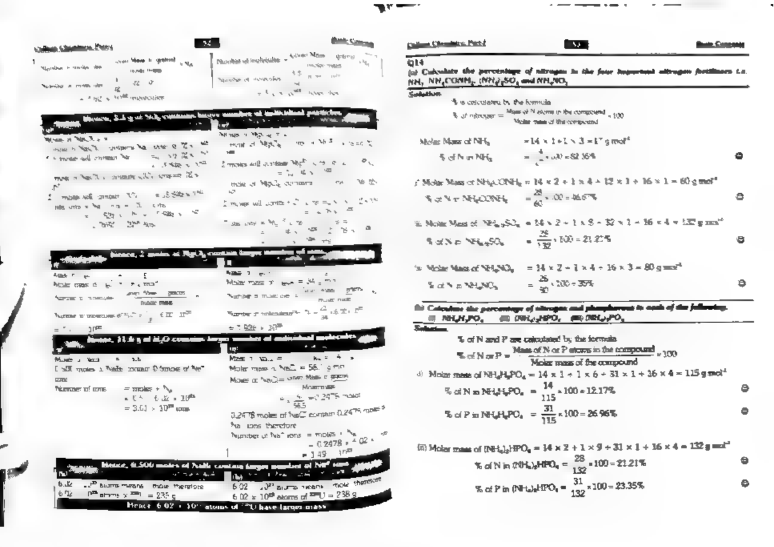
Number of motes = 0.6

Putting the values

Man of Calla

Mass = moles × moler mass

Molar mass of $C_2H_4 = 28 \text{ g mol}^{-1}$



(B) Malar mass of (NH_a)_aPO₄ = 14 \times 3 + $\frac{1}{1}$ \times 12 + 31 \times 1 + 16 \times 4 = 149 g mol⁻¹ % of N in (NH_e)₂PO₄ = $\frac{42}{149} \times 100 = 28.19\%$

% of P in
$$(NH_4)_a PO_4 = \frac{31}{149} \times 100 = 20.80\%$$

Q15. Glucose C_iH_{ii}O₄ is the most important in the cell for generating chemical potential energy. Calculate the mass % of each element in glucose and determine the number of C_iH and O stone in 10.5g of the sample.

Moles mass of $C_aH_{16}O_6 = 12 \times 6 + 1 \times 12 + 16 \times 6 = 180 \text{ g f mos}$

% of hydrogen =
$$\frac{12}{180} \times 100 = 6.67\%$$

% of cosygen = $\frac{96}{180} \times 100 \times 53.33\%$

Mann of
$$C_0H_{12}O_0$$
 = 10.5 g
Miclecules of $C_0H_{12}O_0$ = $\frac{Given'mass (m grams)}{Molar mass} \times N_A$
= $\frac{10.5}{180q} \times 6.02 \times 10^{23}$
= 3.5×10^{23} molecules

$$\begin{array}{ll} \text{1-molecule of $C_0H_{12}O_0$ contains C atoms} &= 6 \text{ atoms} \\ 3.5\times 10^{66} \text{ of $C_0H_{12}O_0$ contain C atoms} &= \frac{6\times 3.5\times 10^{66}}{4.1\times 10^{66}} \\ &= \frac{2.1\times 10^{66}}{4.1\times 10^{66}} \\ \end{array}$$

1 molecule of
$$C_aH_{1d}O_a$$
 contain H atoms = 12 atoms = $12 \times 3.5 \times 10^{48}$ of $C_aH_{1d}O_a$ contain H atoms = $12 \times 3.5 \times 10^{42}$ = $(4.2 \times 10^{49} \text{ atoms})$

$$\begin{array}{ll} 1 \; \text{molecular of $C_aH_{12}O_a$ contain O atoms} & = 6 \; \text{storms} \\ 3.5 \times 10^{44} \; \text{of $C_aH_{12}O_a$ contain O atoms} & = 6 \times 3.5 \times 10^{28} \\ & = 2.1 \times 10^{24} \; \text{atoms} \end{array}$$

College Chemistre: Part-l

55

Q16 Ethylene glycol is used in astromobile antifreces it has 38 7% carbon 9 7% hydrogen and 51.6% oxygen. Its moler mass is 62 ig mot. Determine its empirical and molecular formula.

Solution:

ELIMINI	-	No of grain	Atomic Kario 4	Luquin at Formula
С	38.7	38.7 12 = 3.225	3.225 3.225 = 1	
н	97	97 1008 = 96	$\frac{9.6}{3.225} = 3$	CH _I O
o	51.6	$\frac{51.6}{16}$ = 3.225	3.225 3.225 = 1	

Thus Empirical Formula of Etylene Glycol is CFI₂O Empirical Formula mass = $(12 \times 1) + (3 \times 1) + (16 \times 1)$, = 31 g mol⁻¹

$$n = \frac{\text{Molecular Mass}}{\text{Emprirical Mass}} = \frac{62.1}{31} = 2$$

$$\begin{aligned} \text{Molecular Formula} &= r \; (\; \text{Empirical Formula} \;) \\ &= 2 \; (\text{CH}_1\text{O}) = C_1\text{H}_4\text{O}_2 \end{aligned}$$

Q17 Serviente (H=176 gmol²), is a compound that conducts nerve impulses in brain and muscles, it contains 68.2% C, 6.86% H, 18,09% N, and 9.08% O, what is its molecular formula?

1	EHMINT	* 5 *	No of group #	Atamic Katlo	Loquical Formula
	С	68.20	$\frac{6820}{12} = 5.7$	$\frac{5.7}{0.56} = 10$	
	н	6.86	6.86 1.008 = 6.8	$\frac{6.8}{0.56} = 12$	
	N	15.09	15.09 14 = 1.07	1.07 0.56 = 2	CmH ₁₅ K ₂ D
	D	9 08	9.08 16 = 0.56	0.56 0.56 = 1	

Empirical Formula = C₁₀ H₁₂ N₂ O

Empirical Formula mass = =(12×10)+ (1×12)+(14×2)+(16×1) == 176 g mol⁻¹

Molecular mass = 176 g mol⁻¹ $\frac{\text{Molecular Mass}}{\text{Emprincal Mass}} = \frac{176}{176} = 1$ Q18. An enknown metal M reacts with 5 to form a compound with a formula M_S, § 3.12g of M reacts with exactly 2.88g of sulpher, what are the names of metal M and the compound M₂S₂? The reaction can be written as 2M + 3S - MgS_a \rightarrow Mass of S in M_BS₁ = 32 x 3 = 96 g According to the given data 2.88g of 5 react with M = 3.12 g 2.88g of 5 react with M = $\frac{3.12g}{2.88g} \times 96g \approx 104g$ Thus according to formula M_2S_0 2 moles of M $_{\odot} = 104\,\mbox{g}$ of M $1 \text{ mole of M} = \frac{104}{2} = 52 \text{ g mol}^{-1}$ Hence Atomic mass of M $= 52 \, \mathrm{g \ mol^{-1}}$ Since 52 g mol 1 is the atomic mass of chromium, Therefore M is chromium and MaSa in CraSa La., chromium sulphide. Q19. The octans present in gosotine burns according to the following equation: $2C_0H_{11}+25O_2 \rightarrow 16CO_2+18H_2O$ (a) How many moise of Oz are needed to react fully with 4 moise of octane? The balanced chemical equation for the chemical reaction is $2C_aH_{16} + 25O_a \longrightarrow 16OO_a + 1BH_aO$ Moles of octane = 4 moles

Compare Collag and On according to belanced chemical equation.

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```
Hence Moles of Og required = 50 moles
(b) Hote many males of CO<sub>2</sub> can form one state of octone?

Moles of octane = 1 male
    Compare Caltin and COs according to balanced chamical equation
                      C<sub>6</sub>H<sub>16</sub>
                                  . CO<sub>x</sub>
                                 # 16 moles
                      2 moles
        Therefore 1 mole : \frac{16}{2} \times 1 = 8 \text{cmoles}
         Hence Moles of CO<sub>2</sub> can be produced = 8 moles
(a) How many makes of water produced by the combustion of 6 makes of actuars?

Moles of actuars ~ 6 makes
    Compare CaHas and HaO according to balanced chemical equation.
                      C<sub>2</sub>H<sub>14</sub>
                                 : H₂O
                      2 moles t 18 moles
         Therefore 6 males = \frac{18}{2} \times 6 = 54 moles
    Hence Moles of H<sub>E</sub>O produced. = 54 moles
                                                                                                       æ
(4) If this reaction is to be used to synthanize 2 mains of CO<sub>2</sub>, how many grount of experience 2.
ere needed? How many grams of octans will be used?
              Moles of CO<sub>2</sub> = 8 moles
     Compare CO<sub>2</sub> and O<sub>2</sub> according to balanced chemical equation.
                       CO<sub>R</sub> 1 O<sub>R</sub>
                       16 moles 1 25 moles
                                        \frac{25}{16} \times 8 = 12.5 \text{ moles}
                       8 moies :
     Hence Moles of O<sub>8</sub> required = 12.5 moles
         And Mass of O_k required = 12.5 \times 32 = 400g
```

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C_aH₁₀

1 O, 2 moles : 25 moles

Therefore 4 moles $r = \frac{25}{2} \times 4 = 50$ moles

CO_B : C_BH₁₀

16 moles 1 2 moles

Therefore 8 moles : $\frac{2}{16} \times 8 = 1$ mole

Hence Moles of octane used = 1 mole

And Mass of octane used = 1 x 114 = 114 g

Q20 Calculate the number of gram of Al₂S₄, which can be prepared by the reaction of 20,00g of Al and 30,00g of sulphur. How much non-limiting resciont to in excess?

The testanged chemical equation for the chemical reaction is

Given mass of Al = 20 g

Number of moles of Ai = $\frac{20}{27}$ = 0.74 moles

Given mass of S = 30 g

Number of moles of S = $\frac{30}{32}$ = 0.9375 moles

Determination of Limiting Reactant

Compare Al and Al₂S₃ according to balanced chemical equation

Al : Al₂S₂ 2 moies : 1 mole

Therefore 0.74 moles : $\frac{1}{2} \times 0.74 = 0.37$ moles

Hence number of moles of the product produced by AI = 0.37 moles.

Compare S and Al₂S₁ according to balanced chemical equation.

S I AlgSa

3 males s 1 moles

0.9375 moles 1 $\frac{1}{3} \times 0.9375 = 0.3125$ moles

Hence number of moles of the Al_pS_a produced by S = 0.3125 moles

 Since 5 produces least number of moles of the product, therefore it is the limiting reactant

Hence Moles of Al₂S₈ produced = 0.3125 moles

Mass of Al₂S₃ produced = 0.3125 x 150 = 46.87 g

College Chemistre: Pers-I

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→ Determination of amount of A₁ left unproceed 2AI + 3S → AI₂S₁

Compare the moles of S and A_i to find the moles of All reacted.

S : Al

3 moies : 2 moles

0.9375 moles: $\frac{2}{3} \times 0.9375 = 0.625$ moles

Moles of Al consumed = 0.625 moles

Moles of Al taken = 0.74 moles

Moles of Al laft unreacted = 0.74 0.625 = 0.315 moles

Mass of Allieft unreacted $= 0.115 \times 27 = 9.105$

Q21 A mixture of two liquids, hydresine N₂H₄ and N₃O₄ are used as fuel in rockets. They produce N_2 and water expours Hote many grows of N_0 gas well be formed by reacting 100g of N_2H_4 and 200g of N_2O_4 .

Ф

 \rightarrow The balanced chemical equation for the chemical reaction is $2N_zH_4+N_2O_a \longrightarrow 3N_z+4H_4O$

Given mass of $N_gH_q = 100 g$

Number of males of $N_1H_4 = \frac{100}{32} = 3.125$ motes

Given Mass of N₂O₄ ≈ 200 g.

Number of moles of $N_g O_4 = \frac{200}{2} = 2.17$ moles

→ Determination of Limiting Regulant

Compare N₂H₄ and N₈ according to balanced chemical equation.

N₂H₄ 1 N₈ 2 moles 1 3 moles

3.125 moles $= 1 = \frac{3}{2} \times 3.125 = 4.69$ moles

Compare NgO4 and Ng according to balanced chemical equation.

N_aO₄ (N_a 1 mole ± 3 moles

Therefore 2.17 moles 1 $\frac{3}{1} \times 2.17 = 6.51$ moles

College Chemistry: Farrel Since N_bH_a produce least number of moles of the product, therefore it is the similing Cellege Chemistre: Pers-I 61 . Roper Concepts reactant. Hence Moles of the product produced = 4.69 moles HELLO' Mr Question here! And Mass of the product produced = 4.69 ×28 = 131 32 g Multiple Choice Questions from PAST PAPERS Q22 Silicon carbide (SiC) is an important ceramic material. It is produced by allowing An ordinary mississcope can measure the size of object upto. (Faterlabod Board, 2007) send (StO₂) to react with carbon at high temperature. SiO₁ + 3C → SiC + 2CO (a) 100 nm (b) 200 nm (c) 400 nm Id Jik no cted with excess of curbon, 51 4kg of StC is produced. What is How many times a haemoglobin molecule is heavier than hydrogen atom (Folianished Bread, 2010) Whete 100kg sand to rethe percentage yield of SIC? at 38000 times (b. 58000 times (c) 68000 times (d. 88000 ames Solution: The balanced chemical equation for the chemical reaction is Which of the following element can exist in monoatomic form? (Moltan Board, 2011) $SiO_x + 3C \longrightarrow SiC + 2CO$ (b. chlonne (c) nitrogen (d. relium (a) oxygen Atom of which of the following element has independent existence (Behaviolpus Book). Mass of SiO_a = 100 kg = 100000 gNumber of moles of SiO₂ $= \frac{100000}{10000} = 1666.67$ moles (a) Fluorine (b) Krypton (c) Oxygen (d Mitrogen A Swedish chemist J. Berzeltus determined (Sorgestie Bound, 2013). Compare SiO₂ and SiC according to balanced chemical equation (a) atomic number (b) alomic mass (c) molecular mass (d1 .amc mess ; 51C SiO₂ Electrometer is also called (Lohore Board, 2007). 1 1 mole 1 mole (a) Voitmeter (b. Avometer (c) Ion Collector d Galvanometer Therefore 1666.67 motes t 1666.67 moles Height of peak in mass spectrum shows: (Lahore Beard, 2008: Multion Board, 2023) tal Number of isotopes (b. Mass number Thus (c) Relative abundance (d), Number of protons Moles of SiC, expected to be produced= 1666.67 moles B. Silver has isotopes, rtaken Board, 2010) Motar mass of SiC=28+12=40 g/moi (a) 9 (b) 16 .d) 1B 9. One a.m.u. is expeal to (cohere Sound, 2011 Mass of SiC, expected to be produced=1666.67×40=66666.8 g (a) $1.661 \times 10^{27} \, \mathrm{kg}$ (b) $1.661 \times 10^{27} \, \mathrm{kg}$ (c) $1.661 \times 10^{24} \, \mathrm{kg}$ d) $1.661 \times 10^{24} \, \mathrm{kg}$ 10 Out of 280 isotopes which occur in nature: the radioactive isotopes are (Remodulated Board, -+ Trus Theoretical yield of SIC= 66666.8 g Actual yield of SiC = 51.4 kg = 51400 g18, 116 (b) 126 (c) 30(d) 40 Percentage yield = Theoretical yield 11 The number of isotopes of tip are (Lehors Board, 2011, Re-Spind Board, 2071 (2013) 0., 9 (c) 11 12. The number of molecules in one gram atom of CO₂ is: Kinghamonic Blood, 2016: 51400 66666.8 ×100 = 77% (b) 6.02×10^{22} a. 6.02 x 10²⁵ (c) 6.27×10^{27} (d 6.02 × 10⁻⁴ 13 Empirical formula of glucose is: (Gummunale board, 2911) (a) CHO (b) C₂H₄O₃ (d) CH₂O (dr CaHigO. 14. The mass of ${\rm CO_2}$ containing 8 grams of oxygen ${\rm (O_2)}$ in grams is. (Gipermania board, 2011) a) 32 (b) 22 (c) 16

Book Concepts

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6.2 intere (Cartel) 15 Who determined the atomic mass? (Nata) - Barrel, 20111 td J.Berzelais (c) Newton (b) John Dalton a. Maxwell 16 The percentage of Nárogen in Ammonia is Obston Board, \$013) (d) 78% (c) 92% (b) 46.67% set 82 35% 17 Motecular mass of CaCOs is: 444 (c) 120 (b) 90 (d) 106 a 100 18 Percentage of oxygen in water is dis nd 2009) (b) 88.89% (c) 8.8% (4) 9.8% a 80% lpur Aloural, 20 19 The number of isotopes of program is. (8) (c) Four (d) Three (b) Two 20 Inotopes are saler atoms of same element with similar chemical properties but differen. all Search, 2009). a. Asome number (b) Atomic volume (c) Atomic weight (d Atomic stress 21. The volume occupied by 4.4g of CO₂ at STP is dispute Societ, 2010. (b) 22.4 dm³ (c) 112 cm² (d) 1 12 dec³ a. 2.24 de-1 Z2 The pressure of vepours maintained in unitation charabet of mass spectrometer during MONTHS ATTEMPTE IN Chargedia Sound, 2010) ay 10⁻¹ iour (d)10.4 sam (b) around 10° top: (c) around 10° top: 23. The number of notopies of Cd is. (Baharatpur Board, 2010). (c) 9 0,6 (b) 2 (d) 11 24. The mass of an oxygen atom is (florelying flores, 2020) a) 2667 × 10⁻²⁰ g (b) 2675 × 10²⁰ g (c) 16 g (d) 32g 25. One of the substance is used to absorb CO₂ gas in combustion analysis which is that NUCSTAINCE (D.G Khan Board, 2010) 50% KOH (b. Al₂O₃ (e) Mg(ClO_{4/2} 26 Percentage of N in NH₃ is (Felerial Board, 2011) $40 \cdot \frac{14}{17} \times 100$ (b) 14 × 100 (d) $\frac{3}{34} \times 100$ (c) $\frac{3}{17} \times 100$ 34 27 In combustion analysis, H_cO vepours are absorbed by: (Lehare Beerl, 2007) la) 50% 80H (5) Mg(CIQ₄)₂ (d) MgCl₂ (c) NaOH 28 Positive tons are called: (Lehers Board, 2009) (d) Hydrated of (a) Cations (b) Anions (c) Motecules 29. The volume occupied by 28 g of N_2 at STP is discrepted found, \$410. (a) 22 414 dm³ (d) 1 12 din³ (b) 2.2414 days (c) 224 14 dm³ 30 How many isotopes are present in passadium? (Sergodie Bosed, 2010) a) Four (b) Pive (c) Sac (d) Seven

				-	, ,			21	CK COACAD		
Answ	Answers to Multiple Choice Questions from Past Papers. " ***										
Q#	Ans	Q#	Ana	Q#	Ans	Q#	Аля	Q#	And		
1	(d)	2	(c)	3	(d)	4	ь	5	(b)		
6	(c)	7	(c)	8	(a)	9	(b)	10	fdi		
11	(c)	12	{a}	13	c)	14	(d)	15	ď		
16	(a)	17	(a)	18	(b)	19	,d,	20	fc)		
21	(a)	22	(c)	23	(c)	24	{a ₁	25	cii		
26	(a)	27	(b)	28	(a)	29	(a)	30	C		

Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK-I

SHORT & LONG QUESTIONS FROM PAST PAPERS

ATOM HISTORY OF ATOM, EVIDENCE FOR AN ATOM, MOLECULE

Short Questions

- (1) Write down the main postulates of Dolton's Atomic Theory grave
- [2] Why atoms cannot be observed by an ordinary optical microscope? (Baselylad Board, 1915) Differentiate between home-atomic and hetero-atomic molecule with example at 6. Khan ed. 2011)

(4) Define the term molecule. Give two examples. (Supremble Board, 2015)

one Questions

(1. Define with example, atomicity, molecular formula, Avogadro's number (Messer 8, 2012)

ION, MOLECULAR ION

- Short Overtime

 (1) Describe briefly the energy changes taking place during formation of positive and the formation of lone with respect to expect the expect to expec negative ions. (D.D. Kine Board, 2009) OR Cyrlain formation of lone with respect to energy changes (Nelson Board, \$1989)
- (2) Formation of uni-negative for it an exothermic process. Justify. (Helem Board, 2007)
- | Compation of uni-negative for it an excenerate process stampy from the content of the content

Coffee Cheetern: Part ?

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RELATIVE ATOMIC MASS

Short Questions

- Define Relative atomic mass and give its S I units (Behavelyov Board, 2010).
 What is the atomic mass uit? Give its value in grants. (Festelahed Board, 2009) Gigle Board, 2011. Surpodka Board, 2009)

ISOTOPES

Short Quantitions

- Who are isotopes? From relative atomic masses of isotopes are determined? (Males). Board, 2008, 2011.
- 2 Why the increpes have some chemical properties? (Multan Board, 2009: Reanaplade Board, 2009).
- (3 What are manoisolopic elements? Glue two examples (Roughtud Board, 2013)
- 14. What is the relative abundance of isotopes? How is it determined? (0.0. Khar Board, 2018)
- Lana Questions What are isotopes? Flow do you deduce the fractional atomic masses of elements from the relative botopic abundance? Give one example, (Fetablood Board, 2004, Laborr Sout.
- (2) Explain relative abundance of sotopes with suitable examples. (Familional Bo

MASS SPECTROMETRY

Short Questions

- 1) What is the function of electrometer in separation of isotopes in mass spectrometer
- (Totalstood Board, 2009: Gujressels Boers, 2008)

 (2) Explain mathematical relationship for m/e of an ion in mass spectrometry (Lahors Board)
- (3) What is mass spectrum? (Lakere Board, 2015)
- (4) No individual Ne atom in the sample of elements has mass of 20.18 amu. Explain Why the Bread 2012 Fateshind Board, 2012-Labore Board, 2014) OR Atomic mass may be unition of fractions, Why? (Salabushus Bound, 2009) Multim Bound, 2008, 2010; D.G. Khan South 2009 Labore Bound, 2008. Sergodhe Sound, 2009)

Long Quantitions

- 1. What is mass spectrometer? How is it used to determine the relative atomic mass (and Board, 2012, 2016 Reneipted Searci, 2009 Labour Board, 2008; Sarphi Barry 2009 D.G. Klein Saard, 2012)
- 2 Discuss the mass spectrometer to do the analysis of sotopes of an element. Mention the roles of electric and magnetic fields to separate the isotopes of an element (Mahan Haran
- Flow the untopies of an elements are separated by mass spectrometer makes Board, 2017

EMPIRICAL FORMULA, MOLECULAR FORMULA, COMBUSTION ANALYSIS Short Questions

Oue the main powes for the determination of molecular formula. Ottober Board, 2019

Coffees Chemistry, Part-I

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Basta Concepts

- (2) Define Molecular and Empirical Formula with examples (Schooleger Beard, 2009) Malten Board, 2009: Letters Board, 2007, 2013.
- (3) Molecular formulo is multiple of empirical formula Explain with example in a state
 Board, 2010: Halten Board, 2009, 2011)
- (4, Differentiate between empirical and molecular formula (Februaria Board 2012 Libore Board) 2012: Surgodha Board. 2012 (se part of long q
- 15 COMBUSTION AMALYSIS
- (6) Write formulas used for the determination of percentage of C & H. Hattalahad Board. 2013) (Saroudhe Board, 2012)
- (7) Write functions of Mg(CiO₄), and KOH in combustion analysis (Labore Board, 2014)

Lang Questions

- Write down the steps required to find the emptrical formula of a compound (Guirer Beard, 2008; Behavelers Beard, 2009; Labore Sound, 2014; Mahan Beard, 2008; Cultemanala B
- (2) What is combustion analysis? How the percentages of various elements present in an organic compound are determined? (0.6 Khan Board, 2008. Bahasalpia Board, 2012, D.G. Khan Board, 2009: Gujrumeda Board. 2006) OR Give a method for the determination of the % of carbon and hydrogen in the given organic compound (Labora Board, 2006
- (3) What is empirical formula? Discuss how empirical formula is determined by combustion tricitists. (Estephend Bourd, 2009; Surgodine Board, 2014) 2010: Revealphed Board, 2009) (Gujranusojo Board, 2605 Multan Board

MOLE, AVOGADRO'S NUMBER, MOLAR VOLUME

Shart Questions

Define grom molecule and gram formula.

(Citaben

- (2 NaCl has 58.5 amu as formula mass and not the molecular mass. Justify? (Melton Bound) 1009: Sargodhe Board, 2009: Rasely
- (3) Define Avogadro's Number Give its numerical value? (Subsampler Sound, 1922: D.G. Name Sound, 2008: Multan Board, 2007)
- (4) Define Auggadro's number. How does it relate to the masses of chemical substances?
- (5) Define molar volume. Give two examples. (B.G. Khim Board, 2009)
- (6) One mole of different gases has different masses but occupies some volume. Why? (D.G. Chan Bound, 2010.
- (7) Define (a) molar volume (b, molecular formula (Nassiphult Board, 2012)

Long Questions

- (1) Define the followings giving one example of each UNALAN Bound, 2009;
- (2) (a) gram atom (b) gram ion
- (3) Define and explain the concept of Male and Avagadra's number with examples.
- (4) Write a note on Avagadro's number distant Sound 2015;

Cotton Completes, Part !

Бb

Bank Creek

STOICHIOMETRY

Short Questions

- [1] Give Itso Moichiormetric cassuraptions. (Buhasalpur Bourd, 1 (Oujnapada Bourd, 2011). (Receipted Bourd, 2013: Labour Scart, 2016). 2010), (Lebore Board, 2000
- State law of conservation of mail: (Buhawelpur Bound, 8619)
- (3) Law of conservation of mass has to be obeyed during statchiometric calculations. Give reason or Comment on 8: (D.G. Rhan Band, 2008; Lahors Board, 2015)
- (4) What a statchiometery and define yield of a chemical reaction? (Februaries Bown, 2007)
- 5: Give the assumptions of stoichiometric calculations. (Feintshad Board, 2004,
- Define stoichiometry and turite down its basic assumptions (Fabrical Board, 2018).
 (Insulant Board, 2008). (Surprise Board, 2018)
- 7) Which kind are to be considered during stoichiometric calculations? (Gujernania Bara)
- 8: Write two commption of stoichiometry (Guranuals Board, 2012, 3014: Labora Board, 2014 April 2007)
- (9) By using a bolanced chemical equation, what type of relationships can be studied? Plant Board SRLI
- 1] State conditions of stoichlometry (Revelpted Seed, 2011)
- 2) What is stoichiometry? Give assumption. Mention two improtos was, which help to perform the stoichiometric calculations. (O4) (Labora Board, 2016 Receipted Board, 2012)

LIMITING REACTANT

Short Questions

- (1) How limiting reactant controls the amount of products formed? (Patielabed Board, 2007)
- (2) List steps involved to identify a limiting reactant (relesiabed Board, 2011: Mutan Board, 2010)
- (3) How the limiting reactant is identified? (Dujumede Beard, 2009): (Sergodia Board, 2013)
- (4) Define limiting reactant, give an example. (Labour Board, \$611)
- (5) What is limiting reactant? Write the steps for identification of a limiting reactant, thinks some, 2009. Transforms Sourd, 2010.
- (6) Define limiting reactions. How LR is identified? (Bargoille Board, 2012)

Long Quantions

- Define limiting reactions. How it controls the amount of products? Give example
- 12 What is limiting reactant. Write the steps to identify a limiting reactant. How it contain the amount of products with example (D.O. Khan Board 2018 And Kentuur Board, 2013)

College Christians, Part-

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Book Concesse

VIELD

₩1: === : --- :

Short Questions

- 1) Actual yield is usually less than theoretical yield Why? (Subscalpur Board, 2011, (Emachy Board, 2012) (Onlymousle Board, 2007) (Outronial Board, 2007) (Outronial Board, 2007) (Makenyland Board, 2018) (Makenyland Board, 2018)
- 2) Why the experimental yield is always sess than theoretical yield? (Falsalabad Board, 2013) (Labore Board, 2009) (Helten Board, 2007). Of them Board, 2010) OR What are the reasons for law actuar yield than the theoretical yield of reactions? Mutum Board, 2000) OR Why theoretical yield is greater than actual yield? (Labore Board, 2013), (Multan Board, 2009)
- (3) How can the efficiency of a chemical reaction be expressed? (Gupunusia Board, 2008: Labore
- (4) Why we calculate %oge yield? (Lahors Board, 2014).
- Distinguish between theoretical and experimental visid (Receipted Board, 2009)

Long Questions

- (1) Define octual yield. Write formula for the calculation of percentage yield. (Labora Board.
- (2) What is the difference between actual yield ad theoretical yield? Why actual yield is less than theoretical yield 104 (Lakers Board, 2012)
- (3) What do you mean by the terms: (1) Isotope (ii) Empirical formula: (iii) Gram atom: iv, Yield of a chemical reaction. (Notice Bourd, 2009)
- (4) Define isotopes (Labora Souré, 2012, Maltin Board, 2009—Ran a part of long question))
 - (5) Define yield. How do we calculate the percentage yield of a chemical reaction? (Guiramente Bound, 2003)

(c) What do you mann by make that the way of the same appearance of the company will No good CO have the same number of electrons protein and neutrons explain with re-val. Answer any Eight parts from the followings:

4. Why electrum introductors to used to see the atomic?

4.) The atomic mosses may be in fraction why? (iii) Why limiting reactions control the emount of product during the chemical reaction?
(iv) What is the function of KOH in combination analysis?
(v) Why relative atomic messes of the electrons are in fractions? (vil. What is the pressure of vapours in the ionization chember of male sp The Private as the parameter of requirements in the contactor consequence of makes of the Private Pri x) What is an electromater? (a) What is set determinate?
 (b) It we elements make 50% of the centh crust?
 (c) Whith five elements make 50% of the center equal number of molecules. Give messon?
 (d) One mole of each N₃, CO₈ and H₂ contain equal number of molecules. Give messon?
 (e) A newes are 8th parts from the tottendage.
 (f) In combisation analysis how the percentage of CO₈ is measured?
 (ii) A contiguous has empirical formula HO and its indiscular mass is 34. What is, its protecular formula? (8) The relative abundance of solopes of B is 18 - 200076and 18 - 79,9924e. What is its relative alloyed from? (b) Colculate the mote of Chlorine alterns in 0.882 g Carticol .

(b) Move many motes are likely in expet mosses of C and Mg?

(b) NoCl has 58.5 a mu as formula mass and not the molecular mass shariffy R?

(b) Calculate the mass is grave of 2.74 moles of KMrO_q?

(b) Calculate the mass is grave of 2.74 moles of KMrO_q? INC. Theoretical yield is rese than action yield why? Session. | Astempt any three quantions) (8 x 3)=24
Q5 not What is limiting reactant? Now our you determine (t. Explain it with a chemical reaction
(b) What is an atom? What is its evidence? (00) (03) (c) Colculate the percentage of N and P in NH₄+PO₄

Q6. (a) Whot is stoichiometry. What are its resumptions?

(b) Define the following with examples. (ii) Grain atom. (02 ma: (iii) Gram formula (c) Most of the elements have fractional atomic masses. Why?
(a) Describe combustion analysis for determination of percentages of C. H and O is as 105 organic compound h) NH, gas can be prepared by heating together two subte. NH,Cl and Ca(OH), if a minima containing .00g of which solid is heaten them obsculate the number of grame of NHs produced $2hH \cdot C + C + C + C + H \cdot r_c \longrightarrow C + C \cdot r_c + 2hH \cdot$ 113 Q4. (a) Cartifulate the number of games of R₂SO₂ and water produced where 14g of ROH are reacted with excess of R₂SC₃ arise calculate the number of indecesses of water produced?

Q5. (a) Define with a septiment (in Relative stormic mass. (a) bottopes. (bit) Inch. (b) Define with a septiment (in Relative stormic mass. (b) bottopes. (bit) Inch. (b) Define with a septiment (in Relative stormic mass. (b) bottopes. (bit) Inch. (b) Define with a septiment in the stormic mass. (c) Define with a septiment (in Relative stormic mass. (b) Define with a septiment (in Relative stormic mass. (b) Define with a septiment (in Relative stormic mass.)

Q5. (c) Define with a septiment (in Relative stormic mass. (ii) Inch. (b) Define with a septiment (in Relative stormic mass.)

Q6. (c) Define with a septiment (in Relative stormic mass.)

Q7. (c) Define with a septiment (in Relative stormic mass.)

Q8. (c) Define with a septiment (in Relative stormic mass.)

Q8. (c) Define with a septiment (in Relative stormic mass.)

Q8. (c) Define with a septiment (in Relative stormic mass.)

Q8. (c) Define with a septiment (in Relative stormic mass.)

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Q8. (c) Define with a septiment (in Relative stormic mass.)

Q8. (c) Define with a septiment (in Relative stormic mass.)

Q8. (c) Define with a septiment (in Relative sto 10.1 9 8 g of H₂SO₂ and discolved to water how many regulative terms, profiles 800s, tregetted changes need partitive changes will be rises in the selection.

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Note: Out of Questions 2,3 and 4, Write any TWENTY TWO(22) short unpwers. While writing histories with question number constitute. (mun #)=44 m number ceretally

dection -

QE Another any Eight parts from the followings.

Private that one attent of Mg is finite as heavy as an atom of carbon.

Leve of content into or mass has to be obeyed during allochtomerine calculations.

Fuplain.

One male of different compounds has different means for hearths seems purpose of molecular. Who

College Chamister, Part J

(IVI Delfavorillate between empirical and molecular formula

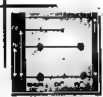
What is the been principle of main spectrometry

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EXPERIMENTAL TECHNIQUES IN CHEMISTRY







Distribution of Is ber Hab and CCle



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Clamin-3

Experimental Techniques in Chemistry

FILTRATION

Filter paper Filter crucibles

CRYSTALLIZATION

Choice of solvent
Preparation of saturated solution
Filtration
Cooling
Collecting the crystals
Drying the crystallized substance
Decolourization of undestrable colours

SUBLIMATION SOLVENT EXTRACTION CHROMATOGRAPHY

Paper chromatography

Objective and short answer, questions (exercise)

Past Papers MCQs and Short Questions

Teat your skills

College Chambers, Part-1

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Exercise and Fachalous in Chamistre



INTORDUCTION

Analytical Chemistre

The branch of chemistry which deels with the study of analytical methods for qualitative and quantitative analysis of chemical substances

Qualitative Analysis

It deals with the Identification of elements, present in a substance.

Occurrentles: Analysis

it deals with the determination of relative emounts of the elements present in a substance.

A complete quantitative analysis involves the following steps

- . Obtaining a sample
- 2. Separation of the desired constituent
- 3. Measurement and calculation of results
- 4 Drawing conclusion

Methods for Separating the Desired Consultant

Following techniques are community used for separating the desired constituent

- A Filtration
- B. Crystallization
- C Sublimation
- D. Solveni Extraction
- E Chromatography

HETRATION

The process of separation of insoluble solid particles from liquids by passing them through a filter medium is called filtration.

- The Insoluble solid particles obtained after filtration is called Residue
- During filtration, the <u>liquid</u> that passes through the filter medium is called <u>Filtrate</u>
- The <u>medium</u> used for fittration is called <u>filter medium</u>

College Chambridge Parks

Experimental Techniques in Charles

Types of Filter Media

The choice of litter media depends upon the nature of precipitate and some other factors

7.4

Two types of filter media are generally used

(1) Filter paper (2) Filter crucibles

(1) Lilla (Popul

A paper used for filtration is called filter paper

Filter papers of different porosities (pore size) are available. The choice of a <u>tilter paper</u> depends upon the <u>size</u> of particles in <u>precipitates</u>.

Whatmann filter paper No. 41 & 42 are in common use.

Nethod of Litration by Class Lennel and Edice Paper

- Filtration by a class funnel and filter paper is usually slow.
- The liquid containing solid particles is poured on a glass rod which then goes into the filter paper
- The residue is collected on filter paper and filtrate collected in the beaker as shown in fig.



englich Indiana as Geble

Palats Line a Good Litration Process

Property of Filter Poper:

(a) The filter paper must be so large that it is ¼ to ½ full of the total precipitate at the end of filtration.

Proportion of Funnel:

- (b) The hannel should be so large that its upper rim is approximately 2_qm above the edge of filter paper.
- (c) The stem of the funnet should be several on long, so that, it goes down into the beaker.

Passention

- (d) To run filtration smoothly, the stem of funnel is <u>kept full</u> of liquid, till there is liquid persent in <u>control part of the funnel</u>
- (e) The lip of the stem of funner should touch the side of beaker, so that the littrate rurs down the side of beaker without <u>solvening</u>.

Cathagret Litter Pages

Following are Important points.

The filter paper should be folded twice

Golfans, Chambeler: Part-C

75 Experimental Techniques in Chemistry

- (a) First fold the paper in half along the diameter of the paper
- (b) Second fold the paper in half again, such that gdgss of the paper do not quite match
- (c) Open the paper on a <u>slightly larger section</u> such that <u>three fold</u> thickness is on one half side and <u>one fold</u> thickness is on other half side and apex englis, slightly <u>greater</u> than <u>60°</u>
- (d) Wet the filter paper with water insert into a 60° degree funnel and firmly press down

Methods to increase the Rate of Filtration

iff Suction Filtentian

The ordinary filtration is very <u>slove</u>. To <u>increase</u> the rate of filtration <u>suction</u> is applied. This is called surfron filtration or vacuum filtration. For <u>better suction filtration</u>, filter paper must be fightly pressed.

(fi) Fluted Filter Paper

- By using fluted filter paper rate of filtration through a conical funnel cast be increased
- To prepare it ordinary filter paper is folded in such a way that a jan like arrangement is obtained
- This fan like arrangement has <u>alternate upg</u> and <u>downs</u>
- Thus, surface area is increased and hence rate of filtration is increased



Control Day Specific

(0) Tar+T = 100

For suction filtration. <u>filter crucibles</u> are commonly used.

<u>Two</u> types of filter crucibles are generally, used.

(a) Gooch Crucible

- It is made up of porcetain baving a perforated bottom
- The perforated bottom is covered with <u>filter paper</u> or with <u>paper unit</u>
- For guick filtration, Gooch crucible is litted into a suction litration, apparatus.
- open Grossmanner
- It is useful for filtration of precipitates, which need to be conited at high temperature.
- Some materials, e.g. conc. HCl. KMnQ₄ solution react with filter paper. Therefore, they
 cannot be filtered using filter paper. For such materials. <u>Assessos Mat is used in place of
 filter paper.</u>



tion Charleton Part I

Experimental Techniques in Chambers

(b) Sintered Glass Crecible

it is a glass crucible, it has a porous sintered class disc sealed into its bottom

It is better then Gooch crucible due to following reasons

- No extra preparation is needed.
- No filter paper is used. Thus, filters of filter paper do not make the product impute.
- It can be gastly handled

Reactive materials (e.g. conc. HCI, KMnO4 solution) can be easily filtered

Difference between filteration through Gooch crucible and Sintered Glass crucible

La made of glass
t has a sintered glass disc sealed into its bottom
Filter paper or Ashestos mal are
not needed
Reactive materials can be littered casily, since no litter paper is used.
Since no litter paper or asbettos mat is used, therefore, no confamination of the product

- ntered glass disc sealed lom er or Ashestos mal are
- naterials can be littered

CRYSTALLIZATION

The process for obtaining crystals of a substance by coaling its but saturated solubles

Crostalization is used to purify crude solid product

Principle

Its principle is that

"Solute is highly soluble in a given joinent at high temperature and excess amount of the solubi comes our as crystais when this solution is cooled."

Mathod

It involves following steps

- (1) Choice of solvent (4, Cooling
 - (2) Preparation of saturated solution (3) Filtration (5) Collecting the crystals

(6: Drying the crystals

(7) Decolourisation of underrable colour

College Chemistry: Post-I

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Exerctor Q9

Write down the main characteristics of a solvent for crystallization of a compo



(1) Choice of Solvent

The solvent is chosen on hit and too basis. Many solvents are tried.

Properties (Characteristics) of an Ideal Solvent.

- (a) it should dissolve a varge amount of solute at its boiling point, and very small amount at room temperature.
- (b) It should not react chemically with the solute
- (c) I should not dissolve impurities or impurities should not gradalize out along with the
- (d) On cooling, it should give well-formed crystals of the pure compound
- (e) It should be <u>cheap</u>
- (f) It should be safe to use
- (g) It should be easily removable

Selvente for Crystaliteation

- Commonly used solvents for crystallization are water rectified spirit (95% ethanol) absolute ethanol, diethyl ether, acetone, chloroform, CCis, acetic acid and petroleum
- If no solvent is found suitable, then a combination of two or more solvents is used.
- If the solvent is <u>inflammable</u> then heating is done on a <u>water bath.</u>

(2) Preparation of Saturated Salation

- The substance is dissolved in minimum amount of the selected solvent
- It is then heated directly or on water both with constant stirring.
- More solvent may be added to the boiling solution until all the solute is dissolved

(d) Libration

- The insoluble impurities are removed, by filtering hot saturated solution
- The solution is littered hot to avoids premature crystallization of the solute on the filter paper or in the stem of funnel.
- Hot water funnel may be used for this purpose.

I' Coding

- Hot beamed notation is cooled at a moderate rate in order to obtain medium seed crystals
- Slow cooking gives bagger stead crystals, which usually contain solvent with impuning Such crystals are difficult to dry.

i Collecting the Cerstals

- When crystallization is complete, the mixities of crystals and mother siquor is hitmed through Grown crucible using a <u>saccum pump</u>
- Full suction is applied in order to remove maximum liquor from the crystals
- If filter cake is band, it is pressed family with a cork to remove remaining mother house.
- Crystals are then washed with small amount of gold solvent several times
- Mother liquor as often concentrated by graporation. It is then cooled to obtain a feet new crop of crystals
- The efficiency of the crystalitization process depends upon the percentage of puring patents obtained from crude substance.

to Decrip the Crystoller d Substance

There are three methods for drying crystals

(i) Pilter Paper:

- Crystals may be dried by pressing between agveral folds of the filter paper. This method
 has two disadvantages.
 - is. The crystals are <u>crushed</u> to a <u>fine powder</u> and
 - (b) The (fitting of filter paper often contaminate the product

(II) Ocean

Drying can be done in an <u>oven</u> if the substance does not <u>mail or decompose</u> on <u>basis</u> (M) Vacuum deslocator:

- Vacuum desector; is a much better method in this, crystels are spread on a watch distance and kept in a vacuum desector for anyeral times.
- The driving agents used in a desertator are CaCl., Silica gol or P.O.

1) Decideorization of Undesirable Colonis

- During preparation of a crude substance, sometimes colouring materials or transitional products.
- Colouring impurities are removed by boiling the substance in the solvent with additional chargost.
- This solution is then littered. The coloured impurities are adsorbed by animal charges
- Pure decoloursed substance crystallines out from the filtrate on cooling.

College Chambrie Park I

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Annual Technique to Change

SUBLIMATION

The process of direct conversion of a solid into repower by heating without penning through liquid phase to called sublimitation.

Solid # Vapour

Sublimation is used to purify soild substances.

The substance obtained after suburnation is called sublimate.

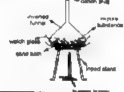
The impure substance, which is sublimed, is called sub-imand

Examples

iodine NH₄Cl. naphthalene, anthracene benzoic acid etc

Mythodi

- The impure substance is taken on a watch glass.
- It is covered with an <u>awarted hone</u>! having a cotton plus into its starr
- The funnel is cooled with wet cotion
- The subscence is <u>breated</u> slowly on a <u>eard bath</u>.
- The pure solid deposits on the inner cold side of the funnal



OFSTRUBUTION LAW OR PARTITION 1 (W.

li stoten

At constant temperature, a solute distributes itself between him immiscible liquids in a constant entits of concentration, independent of the amount of solute added.

Applications

This law is helpful in separation and purification of substances hom mixtures. Two important techniques are based upon distribution law.

10 Solvent extraction (III) Partition Chromolography

Distribution Coafficient

At constant temperature, the ratio of concentration of a substance in two immistible liquids, present to equilibrium with each other, to called destribution on-afficient

It is denoted by K

Mathematically

K - Concentration of a susbance in organic phase is g CCI₂

Concentration of a substance in aqueous phase water)

gen CCl, and water contributing KI er bete he Distribution of local

- Consider the distribution of Is between two immiscible liquids, CCIs and water to the presence of KI
- lodine is insoluble in water. When KI is added in water, it ionizes as
- Jodine rancts with judicle ion to give in-judicle ion in a reversible

l2 + 1° ≠ l2°

Thus, redine dissolves as Is ion in water (aqueous phase).

- Now if CCL, is e-feen to an eq. solution of tri-rodule ion. Lodine is more soluble in CCL. So it moves from aqueous laver to the CCL laver (organic laver)
- As a result, brown colour of to application in acrosous layer factes while the purple color of free sodine appears in CCI tover
- This system of CCL, and H_0O is shaken to increase the area of contact between in layers. Thus, more and more indine moves from equeous to organic layer
- After sometimes, equilibrium is established between the two layers. At this point, thens of movement of I₃ from H₂O to CO₄ becomes <u>equal</u> to the rate of movement of I₃ has CCL to H₂O
- Thus at equilibrium, the ratio of cone of \mathfrak{l}_g in both layers will be constant at constant temperature. This constant is called distribution co-efficient denoted by K and is given in

$$K = \frac{\ln(CG\omega)}{(\xi_2 \cos(\xi_1) \log)}$$

SOLVENT EXTRACTION

The process of obtaining a substance from a solution with the help of an immissibility ent to called poleent extraction.

Principle

- Solvent extraction is based upon Distribution Law
- & solute can be reparated from a solution by an immiscible solvent. The desired solution more soluble in solvent than solution.
- Thus, solute will move from solution to the added solvent layer. This layer can be appropried. separated. The solute can be obtained by evaporating the solvent

Example: Ether extraction, a commonly used laboratory solvent extraction.

It is mainly used to separate organic compounds from water

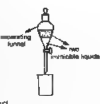
 The aqueous solution containing organic compound is shaken up with ether in a separating funnel. reperating funnel.

College Champions Part-

The organic compound will move from water layer to the ether layer. The morganic impurities remain in the



- Ether is then evaporated to get the pure organic compound
- Repeated extractions using smaller portions of solvent are more effective than using single extraction with larger volume of solvent.
- This technique is particularly useful to obtain the product which is volatile or thermally unstable. Such products are difficult to obtain by other techniques



CHROMATOGRAPHY

It is an analytical technique used for the separation of a minimum, due to differe distribution of substances, between a stationary phase and a mabile phase

e1

Ormin of more

Chromatography is derived from Greek word "Khromaton" meaning "Colour serting"

Stationary Phase

It may be a <u>solid</u> or a <u>liquid</u> supported on an <u>ment solid</u>. Examples: Slica gel, water adsorbed in paper etc.

It may be a <u>liquid</u> or a <u>gas</u>. It flows over the stationary phase Examples, Ethanol, Acetone Hexane etc.

Principle

- It is based upon distribution law.
- The mixture is allowed to come in contact with two phases, a stationary phase and a mobile phase. Different components have different affinities for the stationary phase and mobile phase due to which they are separated.
- The <u>distribution</u> of the component between two phases is controlled by <u>distribution</u> spefficient K given as

K = conc. of a component in the mobile phase cond of a component in the stationary phase Hence, Corepound with amales K value remains with stationers, phoses While. Compound with james K value opes with the mobile phase

Treasure Chambatanaph.

Chromatography is divided into many classes. Two important classes of chromatograph

- 1 Adsorption chromatography
- 2 Partition chromatography

portation Chromotography

In this chromatography, stationary phase is a colld.

When mobile phase flows over the stationary phase, substances staye the mobile than and adsorbed on the stationary phase.

Partition Chromotography

In partition chromatography, stationery phase is a liquid, exported on an inertable in this a substance distributes it self between the mobile phase and the sintionary phase For each class of chromatography many methods can be used

e.g. partition chromatography can be performed by following muthod

Paper Chromatography

	Distance pattern Adeciption and Partition Chromatography							
	Admirentance his narrogeoptic		Proxition a torography upday					
1 2	In adsorption chromatography, stationary please is solid. In this chromatography, solute particles are adsorbed on the	1 2	in partition chromatography- atmonery phase is a fiquid supported on an inert solid. In this chromatography, social distributes shall between					
ı	eolid elationary phase, during their reparation. Example	H	stationary and mobile phase					
1	Thin layer chromelography (T.L.C)	3	Example Paper chromatography					

Paper Chromatography

It is a method of martition chromatography

It has

Sectionary Phone: A liquid adsorbed on a paner

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Mobile Phase A liquid passing over the adsorbed liquid in paper it is issually an ostanic llouid e.g. Ethanol, acetone etc.,

Paper chromatography can be done in many ways

(ii) Ascending (M) Descending (W) Radia/Circular

<u>Ascending</u> paper chromatography is more common.

Assembling Paper Our mentographic

In this method, solvent is placed at the bottom of a vessel. A paper suspended in II. The solvent moves upward by capillary action.

Procedure

- A solvent musture is placed in a chromatographic tank.
- The tank is covered with a lid so that its oner space is judgrated with (Olvent vapours and become homogenous)
- About 20 cm strip of Whatmann's Chromatographic paper No. 1 is
- A line is drawn with a thin pencil, about 2.5 cm above from one end of paper. This is the bose line
- A drop of mixture is placed on the base sine. To identify components, spots of known compounds may also be placed alongside.
- The spots are dried
- The paper is suspended in the chromatographic tank in such a way that the base line must be above the sevel of solvent and the paper is dipped to it depth of 5-6mm. This errangement is left for sufficient time
- When solvens front has moved to ³/₄th of the length of the paper paper is removed from the tank
- Solvent from is marked with a pencil and paper is deted

Chromatogram and its development

- The pattern of spots on the <u>dried paper</u> is called <u>chromatogram</u>
- Development
 - (i) The spots of separated substances can be seen on paper if the eubetances are coloured.
 - (iii) Otherwise, physical or chemical methods are applied on paper to identify apoly of substances



specificace homeonipals.

Each composite of a minture has specific value of Retardation Factor or Retarda-Factor's R_e (subsect.).

or transfed by a component from here here ades is defined as the ratio of the distance is a distance transfed by automs from home line.

R = Distance inveiled by a component from base live Distance travelled by solvent from base line

e.g. From the fig.

$$R_{\epsilon}(A) = \frac{\pi}{\epsilon}$$
 and $R_{\epsilon}(B) = \frac{y}{\epsilon}$

R, value has no unit since it is the ratio of two similar quantities.

- it is used for the <u>mountains</u> and <u>purification</u> of substances from a mixture
- m. It is used for qualitative and quantitative analysis.
- a. It is used to determine the purity of a substance.

College Chambers: Part-I

N1.......

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error Touris

OBJECTIVE AND SHORT ANSWER, QUESTIONS, LIGHTSON

- Multiple Choice Questions.
- (i) A filtration process could be very time consuming if it were not aided by a gentle suction which is developed.
 - a) If the paper covers the funnel up to its circumference
 - (b) if the paper has got small sized poves in it.
 - (c) if the stem of the funnel is large, so that it dips into the filtrate.
 - (d) if the paper fits tightly
- (8) During the process of crystallization, the hot saturated solution; (Secretarial Secret, 8012)
 - (a) is cooled very slowly to g * large sized crystals.
 - (b) is cooled at a moderate rate to get medium stand crystals
 - (c) is evaporated to get the crystals of the product.
 - (d) is mixed with an immiscible liquid to get the pure crystals of the product
- (iii) Solvent extraction is an equilibrium process and it is controlled by
 - (a) law of mass action (b) the amount of solvent used
 - (c) distribution law (d) the amount of solute

(Full-blood Sound, 2007, 2009) (Bahasalpur Board, 2009) (Singodha Board, 2009) (D.G. Khen Board, 2009) (Mallan Board, 2011) (Linham Board, 2010) (Rossiphind Board, 2011) (D.G. Khen Board, 2012)

- (Iv) Solvent extraction method is a particularly useful technique for separation when the product to be separated is:
 - (a) non-volatile or thermally unstable
 - (b), volatile or thermally stable
 - (c) non-volatile or thermally stable
 - (d) volatile or thermally unstable
- (D.G. Khan Board, 2012, (Lahore Board, 2013: Outremedia Board, 2013)
- (v) The comparative rates at which the solutes move in paper chromatography, depend on:
 - (a) the size of paper used.
 - (b) R_i values of solutes
 - c) temperature of the experiment.

(d) size of the chromalographic tank used.
(Behaustur Board, 2008) (Resolute Board, 2009) (Multim Board, 2010) (Surgodha Board, 2018) (Culrements Board, 2018) (Culrements Board, 2018)

ANSWERS TO MULTIPLE CHOICE QUESTIONS it Ans; (d) nit Aust (b) If the filter paper does not fit tightly, suction Slow cooking gives biggger stead crystals, which usually contain solvent with impurities. While will not be developed. fast cooling gives smaller size crystals. Transform, hot saturated solution is cooled at moderate rate to get medium size crystals.

Bear Chambers, Parts!	86	Colin	Chambers Pa
In solvent extraction solute a distributional management in a management of a controlled by distribution law it is AL constant T a solute distribution that it is two immissionals in a constant independent of the empount of solute added	required.		Why is then (D.G. Khon & When a solid rities. To obtain
3 m (b)		901VB	
In paper chromatography the solute move	et according to their K _i values.		
It is defined as R: ** Description of the velled by a co	empowers from base line	Q.5	A water h saltcylic oc separate th
(ii) During filtration, that tip of the beaker to avoid	n is required to dissolve	produ	pitated. Then uct is then cry acid and wat A solid org preparation fover
its boiling point and all [v] Repeated solution extractions tuning a single extraction with its	using small portions of solvent are	condi	he organic co
Answers: (i) qualitative and quantitative analysis inition rease the rate (i.e. large amount.) 3. Tick the correct tentences. If	the sestence is incorrect, write the cored	immis with a sep	ince given on scible with H _e l chloroform in s parated from ound.
GCGCGENENTS.	the manufacture of the second	_	
(a) If the process of filtration is to	run amoubly, the start of the funnel should remain	Q.7	The follows
Correct Statement II the pro	sees of filtration is to run smoothly, the stem of the		Unknown m (NI) Sample (V) Sample

Correct Statement if none of the solvents is found suitable for crystalization

A solute distributes treat between two immucible liquids in a constant ratio of concentrations desired.

Correct Statement A solute distributes that, between two immiscible liquids [1.4] Constant rate, of constant rate, or co

constant ratio of concentrations independent of the amount of solute added Paper chromatography is a technique of partition chromatography

e a need to crystalitie the crude product? cord, 2007: Faisclabad Boara, 2007: Guiranwole Board, 2010: Lehere Scard, 2014) substance is produced during a chemical reaction. If always contains ain pure solid compound, crude product is crystallized from a suitable necluble organic compound septrin is prepared by the reaction of cid with a mixture of acette acid and acette unhydride. How will you a product from the reaction mixture? ation of aspirin, when reaction mixture is thrown in $H_2\mathrm{O}$ aspirin is it is separated from the reaction mixture by suction filtration. The crude stallised from a mixed solvent (i.e., solvent containing equal volume of teri. unte compound is soluble in water as well as in chloroform. During its i, it remains in equeum layer. Describe a method to obtain it from this mpound can be obtained by solvent extraction technique. In the given inic compound is present in squeous layer same a motoroform and chloroform is almost came is bounded and chloroform as O. Hence, organic compound can be separated by shaking aqueous layer a separating funnel. The compound goes into the chloroform layer, which aqueous layer. Then chloroform is evaporated to get pure organic ng Agure shows a deceloped chromatogram on paper with five spots Ixture X (II) Sample A (to) Sample C ₿ (v) Sample D

£1

fet True

(III False Bibl False hi False

CEPTAL SALE

Find out

(f) The composition of unknown missure X

component B and C

The chromatogram shows that the mixture X contains

(10) Which sample is impure and what is its composition

The impure sample is D as it contains A and C.

1111

to solvent extraction technique why repeated extraction using small particular part and more afficient then using a single extraction but larger volume of (Sargedha Board, 2010: Lahore Board, 2011: Rousepind Board, 1981

Repeated extraction using small portions of solvent are more efficient because more emount of substance is extracted.

Write down the main pharecteristics of a solvent for orgentilization of a compound.

Solved on Page 77

Q 20 You have been provided with a estature confutning three inke with different colours. Write dozen the processine to experent the mixture with the help of page chromatagraphy.

into can be separated by following mathod

- A solvent mbdure is pleced in a chromatographic tank.
- The tank is covered with a Rd so that it's inner space is saturated with solvent vapor and become homogenous
- About 20 cm strip of What mann's Chromatographic paper No. 1 is taken
- A line is drawn with a thin pencil, about 2.5 cm above form one and of paper. This is in
- A drop of monare of miss is placed on the base line.
- After drying the spots, the paper is suspended in the chromatographic lank and lehit pullicant time.
- When solvent front has moved to With of the length of the paper, paper is removed to? the tank
- Solvent from is marked with a pencil line and paper is dired.
- The spots of inits can be seen on paper

For each type of talk R, values can be calculated by using the formula

R - Distance treveled by a particular sak from base line Distance travelled by the solvent from base line

ЬЯ HELLO! Mr. Question here!

Multiple Choice Questions from PAST PAPERS

Gooch crucible is made of: (Labora Bread, 2014)

Corner Chemistry, Party

a) Clay (b) Asbestos (c) Porcelain

2 Solvent extraction is a process "Lation Board, 2014)

a) Exothermic (b) Endothermic (c) Equilibrium

(d) Non equilibrium

3. Direct conversion of the solid into vapours is called (Gapus and, 2008, 20181 e) crystallization (b, sublimation (c) distribution

Which one of the following substances is used as decolourizing agent? (Department Board, 100 2009 Mailton Board, 2013)

a) asbestos (b) conc H₂SO₄ (c) animal charoal

5. Substance that does not show the process of sublimation is: (Oak armete Search, 1914, a) K_ICr₂O₁ (b) todine (c) naphthalene (d) NH₄C /

Insoluble particles can be separate from liquid by (renter Bount, 1001)

a) Sublimation (b) Solvent extraction (c) Filtration (d) Crystallization

The drying agent used in a desicostor is (Receipted Board, 2012)

a) NeCL (b) KBr (c) CaCl₄ (d) BaCla

8. Which of the following is purified by sublimation. (Labora Beard, 2009)

(a) Naphthasene (b) Benzoic acid (c) Ammontum Chloride .d) all of these

adine dissolve in water in the presence of 12 due to formation of which one of the following species? (Personal Brand, 2014)

(b) I (c) 1₃*

10. That chemical analysis in which all the elements present in a compound identified is Called (Katalahari Board, 2026)

(a) Quantitive analysis. (b. Qualitative analysis

(c) Gravimetric analysis (d) None of these

11 When hot seturated solution is cooled very rapidly we get. (to

le, medium size crystals (b) large size crystals

(c) premature crystalization of the substance (d) no crystallization

12. The most common solvent used in solvent extraction in procedure and a set of

(a) Acetone (b) Ethanol (c) Rectified spirit (d) Diethyl ether

13 Chromatography is which the stationary phase is liquid is called: (Labor Grand Mer)

(b) partition chromatography lev adsorption chromotography

(d) none of these (c) column chromatography

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Ensuring and Techniques to Charles

14 Which of the following substance shown the property of sublimation? (Surgodia Board id) CH₈COOH (c) Cu5O4

(b) NH₄CI (a) NaCl 15. in chromatography the stationary phase (D.G. Khan Bourd, 2010)

(b) is a liquid a; is a solid

(d. may be solid or liquid c) may be liquid or gas

16. The substance used as drying agent in desiccotor is (Bah 10109 James R. v. (c) 50% KOH (d) Ether b) conc. H₂5O₄

17 A component having small value of K (distribution coefficient) mostly remains in the (a) Akohoi (Supethe Board, 2013) (b) mobile phase (c) chromatographic tank

a) stationary phase 18. Chromatography in which stationary phase is solid is called (Sorgodha Board, 2014)

(a) partition chromatography

(b) thin layer chomnatography

(c) adsorption chromalography

(d) paper chromatography

Austr	ers to l	Multipl	e Choi	ce Qu	stions	from I	ast Pa	pers.	day.
Q#	Ana	0#	Ans	Q#	Ans	Q#	Ane	Q#	Ans
1	(a)	2	(c)	3	(b)	4	(c)	5	(a)
6	(2)	7	(c)	8	(d)	9	(c)	10	(b)
11	(d)	12	(d)	13	(b)	14	(P)	15	(d)
16	(b)	17	(a)	18	(c)		4 30	ang galagas	12

Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK-

SHORT QUESTIONS FROM PAST PAPERS: INO LONG QUESTION

QUALITATIVE QUANTIFATIVE ANALYSIS

Differentiate between qualitative and quantitative analysis. (Multan Board, 2011)

Give the major steps for the complete quantitative analysis, (Mutten Board, 2013)

Nome different methods for separation and purification of a compound (D.G. Khen Board 2008) QR Mention various experimental techniques which are used for the purification of substances. of substances (Lahers Board, 2007)

FILTRATION

What do you mean by (a) chromotogram, (b) filtrate (Resetpinet Board, \$612) (1)

What is filter medium. Name two. (Rasslylad) Board, 2012. (3)

What is meant by filtration and filtration medium (Meter Board, 2012)

College Chambelms Fortel

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- Media used for filtration should be selected on the basis of precipitates. Exploin.
- What should be the site of stem of the funnel used for filtration (Buhavagna B
- How we can run the process of filtration smoothly (Familebad Board, 2009)
- What is fluted filter paper? Give advantage of its use. (D.G. Nhon Bourd, 2011)
- Define filtration. How fluted filter paper is prepared from ordinary filler paper?
- dha Board, 2810)
- How does Gooch crucible increases the rate of filtration? (Surpodia Board, 2009) OR Expirely juiration through Gooch crucible (Resetpinet Board, 2011
- 10) Differentiate between filtration by Goods crucibie and Sintered glass trucible (0.6)
- Why sintered giens crucible is preferred to Gooch Crucible? (Musican Bound, 2012)
- Concentration HCl and KMnO₄ solutions cannot be filtered by using filter paper Discuss (Gujrenucle Board, 2011 Behausiper Board, 2011 Februlaburé Board, 2017) OR Hou can HCl and KMnO4 solutions can be filtered by Gooch crucible? (Bargotha Board, 2012)

CRYSTALLIZATION

- What is (or Define) crystallization? (Fabrithed Board, 2008: Gafrenucia Board, 2008: Surgette
- Mention the major steps involved in crystallization. (Revolpted Board, 2007, Labora Bo
- Give the main characteristics of the solvent used for crystallization (Lohors Bourd 2009; Fotsalabod Board, 2009; Fotsalabad Bourd, 2019; Multon Bourd, 2019, Gulymounte Bourd, 2019; D.G Khan Board, 2012; D.G. Khen Board, 2012; Azad Kashmir Board, 2012; Bahmaelyur Bourd, 2012; Gujranusia Board, 2012) OR Give the sollent features of an ideal solvent (Surgodia Sound. 1015) (NOTE: Sometimes two or four characteristics are asked)
- Name jour (or eight) important solvents chosen for crystallization. (Petelahad Board, 200) Repolpinal Board, 2011. Labore Board, 2013;
- What is disadvantage of slow cooling in crystallization? (Fetochhed Bours, 2013)
- Describe method to collect crystals from mother liquor (Sergodia Bound, 2011)
- How vacuum desiccotor is used to dry the crystals (Matter Board, 2008, Bake 2009. Sargodka Board. 2013) OR The desiccator is safe and reliable method for drying the crystals Expinin (Lahore Board, 2012)
- Name two drying agents used in vacuum desicozion (D.G. NAMI Bourd, 2009) OR Name the chemicals which are used as dryig agents in the desiccotor? (Muten Board, \$011)
- How can you remove undestrable colour from the crystals (Huban Bound, 2007 A Starrd, 2007: Bahmsalpur Bourd, 2008: Gujtonisale Board, 2008: Emesiphed Sourd, 2010; OR House the colouring impurities are removed from a crude substance? (Recognical Board, 2013)

SUBLIMATION

- (1) Define sublimation with examples OR What is sublimation? Name any two sublimates Of Sublime Sciids) (Lakere Board, 2008: Rauslohell Board, 2010: D.G. Nhan Board, 2012: Gefrancele Board, 2013: Lakere Board, 2014: Makan Board, 2008: Bakewelper Board, 2011: Ge Board, 2024. Sergodho Board, 2019)
- Define sublimand and sublimate (Labora Board, \$014)

Esperimental Techniques in Chambring

- Expirin the process of sublimation. (Falmished Board, 2012)
- How nophtherene can be purified? (Gapmanh Board, 2008; Recompted Board, 2009) Of Naphthalens can be best purified by sublimation. Why? (Raudphait Board, 2006) (4)
- Give the importance of sublimation. (Labora Beard, 2013)
- Why solid incline sublimes? give reason (Surgodia Bourd, 2007)
- Differentiate between sublimation and condensation (Muham Boord, 2009) (7)

SOLVENT EXTRACTION, DISTRIBUTION LAW

- What is solvent extraction? (Aund Keshati Board, 2019: Labore Board, 2008, 2013) OR Side solvent extraction and give importance. (Sergethe Board, 20) t
- Define distribution law (or Partition law) and how it is helpful in solvent extraction?

 Labora Road, 2010; D.G. Khan Board, 2011; Gajanneda Board, 2011; Sargodhe Board, 2018; Bakonalper Board, 2010; B.G. Khan Board, 2018; Gafranteda Board, 2014;
- What is distribution coefficient? To which technique it is applicable. (Gapuates): D.G. Khan Board, 2016)
- lodine is more soluble in water in presence of KI. Discuss. (Februaries at Bound, 2011)

CHROMATOGRAPHY

- Define chromatography and give formula of distribution coefficient. (Gapanusis Book **2011**)
- What is chromatography? Give its two types. (D.G. Khas Board, 2007)
- Define chromatography and explain adsorption chromatography. (Netter Booms, 2016)
- OR What is adsorption chromatorgraphy? (Bahanatan Board, 20).
- Define/Differentiate between adsorption and partition chromatography (D.G. (Non-Bert 2009: Lehrer Board, 2011: Malten Board, 2019: Serpodue Board, 2013, 2014: Lehrer Board, 2019) Define (a) partition law (or Distribution law) (b) chromatography (Malten Board, 2013)
- (6)
- 3011) (8)
- Differentiate between stationary and mobile phase (Notice Board, 2009: Feb. rale Board, 2014)
- chromatograph/ What do you know about R, value of a component in paper (9) (Deplements Source, 2015) OR What is R₁ value? Give its formula, the stan Board, 2007 of What is R, value? Why it has no units? (Lakare Board, 2009; Bahmelipin Board, 500). February 2010; February 2017; Western Board, 2012)
- Give two applications of paper chromatography (Betweetper Board, 2012)
- (11) Define chromatography. Give its uses 10-1-manls Board, 2009: Labore Board, 2010, 2016 ndi Board, 2013)
- (12) R, value is always less than 1.0. Comment on it. (Patentaled Board, 2008)
- (13) What is the role of stationary phase in chromatography? (Surgedia Board, 2007)

College Cheroleten Part-

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Experimental Techniques in Chemistre

1151 VOUR SKILLS Marks 85 ORIECTIVE Marke: 17 Time: 20 Moutes ver writing, cutting, estating, using lead penalt will result in loss of marks Q1 Each question has four peerible answers. Choose the correct answer and encivels if (i) The choose of filer media depends upon (a) ontitue of reactants. (b) nature of reaction (c) nature of Precipitates. (d) nature of filer paper Filtration by a glass functor and filter paper is very (a) slow (b) difficult (c) fast (d) acc. Goods crucible is not used to filter the solution of (d) accurage [III] is glucoss (b) RMnO c) KOH id K_SO, in crystallization, solvent should dispolve more solute at (a) freezing temperature (b) room Temp (c) high temperature (d) low temperature (b) room Temperature (c) rapp remperature to low importation in solution by using the solvent or which the solute is (a) more soluble. (b) portially soluble (c) insoluble. (d) soluble at high remperature. Repeated extraction using small portions of solvent one more (a) occurate (b) efficient (c) slow (d) regard Solvent extraction is based upon (a) mass law (b) distribution raw (c) avegadre s few Which of the r oblawing a used as drying agent in vacuum decisioner? Which of the following is used as drying agent in section descriptor? Lighting pell b) Phosphorus inchered a (2 lactum sulphake d) None A component having small value of K (distribution coefficient) mostly etimisms with Ley Stationary phose (b) Mobile phase is Chromatographic tank Which of the following substance shows the property of subtrination? A) Sodiam chloride [5] announced chloride (2 Copper sulphate d) acetic Which separating techniques at based upon Distribution Law? A) crystalization b) subtimation (b) tolored extraction (d) fibration Blue of Structure can be increased upon id) None of these c) Copper sulphate d) acetic acid (m) Rate of Eltration can be increased using (b) Chromatographic tank (c) Loid finger (a) Detacation (d) Suction Bask A modute of tries is separated by paper chromatography due to b) the different R₄ values of the Iniq. the amount of solvent taken. d) amount of tales use c) the temperature of the system in which technique a solute distributes between two imprescible impude? a) crystallization b) solvent extraction c) Fiftion d) di a) cystalization (b) sobject extraction. The safe method of drying crystals is by using a) oven b) vector desicator c) folds of filter paper d) surflight Which one of the following substances is used as decolorating agent? a) eclassics b) cone H_aSCl_a d animal charces d) which get Paper chromatography is a feeting a col a) adsorption chromatography b) partition thromatography c) solvent chromatography d) both (a) and (b) SUMMECTIVE Market 60

Times: 2:10 House Maste: Dut of Questions 2,3 and 4, Write any TWENTY TWO(22) short on Northern carefully. (22: 21=44 Section.

(ii) Suitered glass crecible at better than Conch crecible Why?



(BI)

(a)

西班西西西

- (iii) Differentials blw edeorption & partition disconsatography.

 (iv) What are different methods used to day crystals?

 (ii) White down the uses of chromatography.

 (vi) What are the major steps for complete quantitative determination of a sample?

 (vii) Yours the solvents commonly used in crystallisation.

9.4

- twin reams the universe commonly used in operations or in the process of crystallization.

 (viii) How the undestrible colour can be removed in the process of crystallization.

 (b) Differentiate between qualitative and quantitative analysis.

 (c) Negree the substances that can be purified by the process of additionation.

 (ai) Write down the distribution equation for the distribution of I₂ biv CO₂ and H₂O?

- (a) White down the distribution equation for the distribution of I₂ over CLL₂ and ri₂O?

 (di) What is analytics chemistry?

 (3) Assert step Eight parts from the followings.

 (i) How does Gooch Cruzible increases rate of filtration?

 (ii) Why fained filter paper is used?

 (iii) Why concentrated CHC and Rolfo, solutions cannot be libered by Gooch Cruzible?

 (iv) White the nauces of the major steps involved in the crystallization?

 (v) What is matter laptor?

- (v) What is mother laquor?

 (a) What is the sale and reliable method of daying crystals?

 (vii) Why crystalization is used?

 (vii) Write down the main characteristics of the solvent used for crystalization?

 (vi) Define distribution Law or partition Law?

 (a) How the methors of nephthalene and wand to separated?

 (d) What is other sotroicalor?

- (iii) What is paper chromotography? What are its types? Q4. Assure any Sta parts from the followings. (ii) How S₆ upbe is micralated?
- £13

- (u) Why hat fitnesson is necessary during crystallization? (uii) What is decommandered? (uiii) Describe method to collect crystals from mother liquor?
- (b) How todine is dissolved in water?

- Section | | (Attempt any three questions) (6x \$)=24
 Q6. (a) Write down five characteristics for an ideal solvent for crystolization?
 - (b. What is the disedvantage of slow cooling in crystallization?
- (c) What is the basic principle of crystallization? Q6. (a) What is distribution law? Give examples.
- (b) Define the following:
- (ii) Subtraction (ii) Chromatography

 Q7.(a) What are the important considerations for a good (Unation process.
- (b) Why repeated admittions using smaller portions of activant are more beneficial than using smaller portions of activant are more beneficial than using smaller portions of activant are more beneficial than using smaller extraction with larger volume of solvent?

 (c) How school extraction is corried out?
- QB.(a) How many different ways are there to carry out paper chromatography? Briefly describe the procedure for according paper chromatograph.

 (b) Stem of funnel in filtration should be several inches long. Why?
- ic) What type of substances cannot be filtered through Gooch crucible using filter paper?
- Q9. (a) What are the common laboratory superimental techniques?
 (b) A solid organic compound is soluble in water as well as in chlorolorm. During its preparate.
 - to squeous layer. Describe a method to obtain it from this layer (b) How will you prepare a fluted filter paper?







Lightening bolts(Pir

*/ P A / 7 TO A



STATES OF MATTER

Chapter-3 Rose . Tweeting home

Properties of gases Properties of liquids Properties of solids Units of pressure GAS LAWS Boyle's law Experimental verification of Boyle's law Graphical explanation of Boyle's law Charles's law Experimental verification of Charles's law Derivation of absolute zero Scales of thermometry GENERAL GAS EQUATION ideal gas constant 'R' Density of an ideal gas AVOGADRO'S LAW

DALTON'S LAW OF PARTIAL PRESSURE

Calculation of partial pressure of a gas Applications of Dalton's law of partial pressures DIFFUSION AND EFFUSION Graham's law of diffusion KINETIC MOLECULAR THEORY OF GASES Explanation of gas laws on the basis of kinetic molecular theory KINETIC INTERPRETATION OF TEMPERATURE LIQUEFACTION OF GASES General principle of liquefaction Methods for liquefaction of gases NON-IDEAL BEHAVIOUR OF GASES
Causes for deviations from deality van der Waal's equation for real gases
PLASMA STATE Objective and short answer, questions (exercise)

Numerical problems (exercise) Past Papers MCQs and Short Questions

Test your skills



STATES OF MATTER

Matter exists in four states.

(I) Gas (II) Liquid ## Solid

(b) Plasma

Liquids are less common than solids, gases and plasmas. It is because the aquid state of any substance can exist only within a relatively narrow range of temperature

Properties of Gases

- 1. Indefinite Volume. Gases have indefinite volume. The volume of the gas is equal to the volume of container
- 2. Indefinite Shope: Gases have indefinite shape. They adopt the shape of the container in which they are stored.
- 3. Low density: Gases have yow densities than solids and liquids. Thus gases bubble through liquids and tend to rise up
- 4. Jours-Thomson Effect: When gases expand suddersly, they cause cooking. This effect is known as Joule Thomson effect. This effect is used in the liquelaction of gases on Industrial scale
- Efficient & Diffusion: Gases can offuse and diffuse.
- Effect of Temperature: Gases expand on heating & contract on cooling tell increase. In temperature increases the volume of gas & vice versa. Liquids and solids do not show considerable increase in volume by increasing temperature
- 7 Effect of Preseure: Gases expand on decreasing pressure and contract on increasing Dressure.
- 8. Gas Pressure: Gas motecules are in constant random motion. During their motion. they collide with the wall of container. These collisions develop a pressure called Gas
- 9. Intermolecular Porces: Gases have very weak intermolecular forces.



Definite column: Liquids have definite volume.

 Schools Shape: Liquids have indefinite shape. They adopt the shape of the container in which they are stored

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- 3 Constant motion: Its figures, motionies are in constant motion. Evaporation and diffusion properties of liquids are due to this motion.
- 4. Demaittee: The demaities of liquids are greater than gases and close to solids
- Intermolecular forces: The intermolecular forces among the liquid molecules are stronger than gases but weaker than solids. The malting points and boiling points a liquids depend upon the strength of these forces.
- 6. Spaces: The spaces among the liquid molecules are negligible just like solids
- 7 Kimstle energy: Liquid molecules have K.E. Liquids can be converted into solids by cooling i.e. by decreasing the K.E. of the molecules.
- 8. Collegions: Molecules of liquids collide with one another and exchange energy
- 9 Diffusion: Liquids can diffuse into other liquids. However, the rate of diffusion is smaller than that of gases

Properties of solids

- I Definite shape: Solid have definite shape.
- 2 Definite solume. Solid have definite volume.
- 3 Spooses Molecules of solids are very close to each other. They are tightly packed.
- 4. Incompressible: Solids are incompressible due to tight packing.
- 5. Intermolecular forces: Solids have strongest Intermolecular forces.
- 6. Vibrational motion: Solid particles only show vibrational motion.
- 7 Differitors: Solids have negligible rate of diffusion.

Pressure and Units of Pressure

Pressure is the force per unit area is $e^{-P} = F \wedge A$

- = The SI unit of pressure is Nm ⁻³ (Pascai). 1Pa=1Nm, ⁻³
- The unit pound per square inch (psi) is most commonly used in engineering work.
- · The unit milibar (mb) is commonly used by materologists.
- in chemical work, pressure a defined in terms of standard almospheric pressure. The pressure of oir hat can support 760 mintig column at sea level, is called one amountable to the force exerted by 760mm or 760m long column of mercury on an area of terms to 390.

It is the average pressure of atmosphere at sea level. Its symbol is 'etm' 50, one atmosphere at sea level. Its symbol is 'etm' 50, one atmosphere at sea level.

Interconnersion of parious pressure units

1 atm = 760 torr = 760 mm of Hg = = 101325 Nm^{-E} = 101325 Pa = 101325 $^{1/2}$ (kilopateal) = 14.7 pound inch $^{-4}$ = 1013.25 millibara.

LASTAICS

All gases behave uniformly. When pressure and temperature are changed, the volume of ones is changed. The gas laws describe this uniform behaviour of gases.

The relationships between volume of gases and the external conditions like semperature and pressure are called the gas laws.

Exercise Q 4. (a)

What is Boyle's law of gases? Gives its experimental verification.



Bogle's Larg

Robert Boyle gave this law in 1662 It states.

At constant temperature, volume of a given more of gas is inversely proportional in the pressure exerted on it.

Mathematically
$$V_0, \frac{1}{p}$$
 (When 'T' and 'n' are constants)

or PV = k _____(1) (When 'T' and 'n' are constants)

Eq (1) gives another statement of Boyle's law

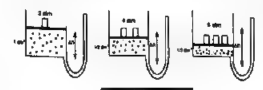
At constant temperature, the product of pressure and volume of a given mass of a gas to always constant,

If the pressure of a given mass of gas is P_1 and volume V_1 , then by changing pressure to P_2 , volume is changed to V_2 , such that

So
$$P_1 V_1 = P_2 V_2$$

The second of the

- Consider a gas in a cylinder fitted with a movable piston at 25°C. The cylinder is fitted with a manipreter to read the pressure of the gas directly.
- Let the initial volume of gas enclosed in cylinder is 2 dns and pressure is 2 atm.



100

Then according to Boyle's taw

$$P_{i}V_{i}=2\times1=2~\text{atm dm}^{2}~\text{where is}$$

When pressure is doubled on the gas, its volume becomes half

Therefore

$$P_aV_a=4\times 16=2 \text{ atm des}^a=k$$

Similarly if pressure is tripled (i.e. 6 atm) on the gas, its volume is reduced to 1/3 Therefore

$$P_aV_a=6\times 1/3=2 \text{ ann dm}^a\equiv k$$

Street product of P and V is always constant. Hence Boyle's law is verified

A gas having a volume of 10 dm^3 is exclosed in a passet at 0°C and the preatmospheres. This gov is allowed to expand until the new presence in Z atmosphism will be the new volume of this gar, if the temperature is maintained at 273 K7.

Selection.

 $V_1 = 10 dm^3$

$$V_2=2$$

 $P_1 = 25 \text{ atm.}$ $P_e = 2 atm$

Since the temperature is constant ($T_1 = T_0$), thus according to Boyle's law

$$P_tV_t = P_tV_t$$

$$V_g = \frac{P_1 V_1}{P_2}$$

$$V_{p} = \frac{2.5 \text{ atm} \times 10 \text{ dm}^{2}}{2 \text{ atm}} = \boxed{12.5 \text{ dm}^{0}}$$

coupl is all explanation of Boxle's for

Exercise Q.4 (b).

What are inotherms? What happens to the positions of teatherms when they are plates (Loncee Board, 2614 high comperature for a particular gas?

Graph betream V and P (leatherms)

- The plot of volume of a gas on Y axis and pressure on X-, axus at constant temperature (e.g. at O^C) gives a curve called trotherm.
- If this graph is plotted at higher constant temperature (e.g., at 25°C), then the curve goes owby from both axis. It is



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because, at high temperature, the <u>volume</u> of gases is <u>increased.</u>

Similarly, if curve is plotted at further higher temperature, then the curve further goes ·/

Emercise O.4 (b)

Why do we get a straight line when pressures exerted on a gas are plotted against inverse of volumes? This stroight line changes its position in the graph by varying the temperature Justify it

Graph between P and 1/V

 The plot of <u>pressure</u> (P) on Y axis against inverse volume <u>1. V) of a</u> gas, on X-axis gives a straight line, it is because increase in pressure decreases the volume hence inverse of volume increases. Thus, 'P' is directly proportional to 1/V



 At higher temperature straight line becomes closer to Y-axis. It is because at higher temperature volume of a gos is top to April 2 two of April 2. Increased, therefore 1 / V is decreased.

The straight line passes through the origin because when the pressure is close to zero then volume is so high that I/V is very close to zero

Graph between PV and P

- A plot of "P" on X-axis against "PV" on Y -axis will give a straight line parallel to X exis. The straight line shows that it constant quantity
- At higher temperature, volume of gas is increased. But new value of PV also remains constant. Thus, a straight line parallel X-axis is again obtained at higher temperature.
- This straight line helps us to understand the non-ideal behaviour of gases.



IS a

Boyle's loss is only applicable to ideal;

Exercise Q5 (a)

What is the Charles's law? Which scale of temperature is used to verify that $\sqrt{T}=\kappa$ (pressure and number of motes are constant)?

CHARLES'S LAW

Chades, a French scientist, dave this law in 1787.

If status

As equalized promotes, the notions of a fourt union of got in directly proportional to ϕ_{ij}

وقد تا المحطورة (

$$\frac{v}{T} = k$$
 (1)

where T is the temperature in Kelvin.

Alno

$$\frac{V_1}{T_1} = \frac{V_2}{T_2} = k$$

Eq. (3) also shows that

At community presence, the ratio between the column of a given mass of a ges and it absolute temperature is always constant.

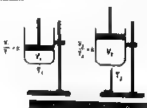
Sec. and

The graph between 'V' of a given mass of ges and 'T' will be a straight line. It is because, 'V' is directly proportional to the absolute temperature.



specimental Verification of Charles of the

- Consider a gas, enclosed in a cylinder fitted with a moveable piston. At temperature To
 the volume of gas is V₁.
- . By increasing the temperature to " T_z ", its volume is increased to " V_z ". The pressure on the piston is kept constant.



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 It is observed that ratio between volume of a given mass of gas and temperature remains constant at constant pressure

$$\frac{V_j}{T} = \frac{V_j}{V_j} = k$$

Artification of Temperature Scale for Charles's Law

The mathematical form of Charles s law is

$$\frac{V}{T} = X$$

This equation only holds if temperature is on Kelvin scale Example:

Consider the following data for a gas when the temperature is on Keivin scale

U disq	1092	846	746
Title 🏢	546	423	373
VT (dm 'K ')	2	2	2

Hence value of $\frac{\mathbf{V}}{\mathbf{T}}$ is always constant

However if temperature is taken on Celaius scale, then the above relation is not satisfied. e.g. Consider the following data for above gas when the temperature is on certigrade scale.

V (dm 1)	1092	846	746
1000	0.	150	100
V1 (dig) *C 1	4	5.64	7.46

Hence, value of $\frac{V}{T}$ does not remains constant.

Thus, Charles a law is only obeyed if temperature is on Kelvin scale

Example 2;

250 and of hydrogen is cooled from 187°C to -27°C by melmolning the pressure constant. Calculate the was volume of the gas at loss temperature.

Plution

$$V_{\rm f}=2$$

$$T_1 = 127^{\circ}C + 273 = 400 \text{ K}$$
 $T_2 = -27^{\circ}C + 273 = 246 \text{ K}$

According to Charles's law

$$\frac{V_1}{T_1} = \frac{V_2}{T_0}$$

$$V_2 = \frac{V_1 \times T_2}{T_1}$$

$$V_s = \frac{250 \times 246}{400} = 153.8 \text{ cm}^3 = \boxed{0.1538 \text{ dyn}^3}$$



Exercise Q.6. (e):

What is Kelvin scale of temperature? Plot a graph for one more of a real gas to prove that a gue becomes liquid, earlier than -273 16°C. (Graphical Explenation)

Absolute Zero

The hypothesical temperature at which the volume of all gases becomes zero is collect

For routine calculations the value of absolute zero is taken as 273°C

Mathematical Lyphanation: Quantitative statement of Charles's Inc.

At constant pressure, the solume of a given mass of an ideal gas increases of secretary 1/273 of its original values at 0°C for every 1°C rise or fall in temperature

Thus Volume at T*C =
$$V_{\tau} = V_0 \left(\frac{273 + T}{273} \right)$$
......(1)

where V_T = Volume at T⁴C , T=Temperature in ⁴C

Therefore at ~273°C

$$V_{sph} = V_0 \left(\frac{273 - 273}{273} \right) = 0$$

- The comparature -273 °C is called Absolute Zero of Kelvin scale. Thus volume of a 950 becomes sero at absolute sero
- Absolute zero can never be achieved, it is considered as the lowest temperature. Its value is independent of the nature of gas.
- Charles a law is not obeyed when the temperature is on Centigrade scale. For this reside Kelvin scale was developed with 0 K = -273 °C

College Chambelra: Part-1

Graphical Leplanation · According to Charles's law, when a graph is plotted between V and T for a gas, a straight

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ine is obtained

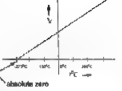
This line intersects the temperature exis at 273°C, which is considered as the lowest temperature

 This temperature is achieved if the substance remains in the gaseous form. But all gases liquely before reaching this temperature.

Thus, the lines of all the gases are extrapolated. They meet T' axis at -273 °C at which the volume of all gases becomes zero. However, it can never happen for a real

gas. Thus, 273 °C is taken as zero of Kelvin scale and it is called absolute zero.

Generally, greater the mass of the gas greater will be the slope of the line. It is because greater number of moles of gas occupies more volume



SCALES OF THERMOMETRY

The branch of ectence which deals with the measurement of temperature is called thermometry.

There are three measuring scales of thermometry

(I) Centrigrade or Calsius Scale (II) Fahrenheit scale

(III) Kelvin Scale

(i) Centigrade or Calaius Scale (°C)

The temperature measured on this scales is represented by "C

On this scale freezing point of water is marked as 0 °C and boiling point as 100 °C. The distance between these two ends is divided into 100 equal parts. Each part is equal to 1°C

(II) Pohrenheit Scale: (*F)

The temperature measured on this scale is represented by "F

On this scale the freezing point of weser is marked as 32 °F and boiling point as 212 F The distance between these two ends is divided into 180 equal parts. Each part is equal to 1°F

(III) Keluin seals (Ki

The imperature measured on this scale is represented by K Interconcersions of different scales of thermometery

Fahrenheit to Celaius

$${}^{\circ}C = \frac{3}{9} ({}^{\circ}P - 32)$$

Celalus to Fahrenheit

$$^{6}F = \frac{9}{5} \, ^{6}C + 32$$

Relvin to Celsius and vice versa K⇒*C + 273.16

100

What is the general gas equation? Derive it in various forms?

GENERAL GAS EUQATION

For a given mass of gas, the three variables i.e., pressure, temperature and volume can be inter-related by one equation import as "General Gas Equation" or "ideal Gas Equation"

According to Boyles law

"At constant temperature, values of a given mass of a gue to inversely proportional to the pressure exacted on it" Le:

$$V_{\text{CL}} \frac{1}{9}$$
 (at fixed T & n) ____ (1)

According to Charles' law

"As constant pressure, the polume of a given mass of a gas is directly proportional to chechate temperature." i.e.

The volume of a gas to directly proportional to the mass (moles) of the gas at stant temperature and pressure 1.8;

Combining eq. (1), (2), (3),

where 'R' is constant known as universal gas constant.

Eq. (4) is known as general gas equation

For 1 mole of a gas

$$PV = RT \quad \text{or} \quad \frac{PV}{T} = R$$

If P,V & T are changed for a gas from $P_1 V_1 \& T_1$ to $P_2 V_2 \& T_3$

Then
$$\frac{P_1V_1}{T_1} = \frac{P_1V_1}{T_2}$$

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Reduction of Content Cas Ego too to Buch shar Charles's law and Accomples Sag.

) T' and 'n' are constant then Pv = k (Boyle's Law)

or
$$V = \frac{nRT}{m}$$

If ${\mathcal P}_{-}$ and in are constant then $V \approx kT$ (Charles 5 Law)

or
$$V = \frac{nRT}{p}$$

if 'P' and 'T' are constant then $V=k\,n$,Avogadro's Law)



Exercise Q7 (b):

Can we determine the molecular mass of an unknown gas if we know the pressure. temperature and solume along with the mass of the gas

Molecula: Mass of Guses

The general gas equation is

Since

Number of moles =
$$n = \frac{Glver mass}{Molecular mass} = \frac{m}{M}$$

So equation (1) becomes

$$PV = \frac{m}{M} \times RT$$
 _____(2)

$$or. M = \frac{mRT}{cst}$$
 (3)

Thus knowing, pressure (P), volume (V), temperature (P) and mass (M) of a gas, its molecular mass (M) can be calculated.

Exercise Q7 (c):

How do you justify from general gas equation that increase in temperature or decrease of Pressure decreases the density of the gas?

1 atm = 101.325 Nm⁻⁸

 $1m^0 = 1000 \text{ dm}^0$

Deposit of Come

A. THE PROPERTY OF PERSON AND PROPERTY OF PERSON AND PROPERTY OF PERSON AND PROPERTY OF PERSON AND PERSON AND

The general gas equation is $PV = nRT ____(1)$

n = Molecular ment M

So eq. 4) becomes

$$PV \circ \frac{m}{M} \times RT \longrightarrow (2)$$

Since,
$$d = \frac{m}{V}$$
 so eq. (3) becomes

- Finouring, pressure (P), temperature (T) and molecular mass (M) of gas, its density (d) can he calculated
- The equation shows that density of an ideal gas is directly proportional to its molecular mass and pressure and inversely proportional to the temperature. So, increase in M' and P', incremes the density while increases T' increases decreases the density due to increase in volume

THE PURPOSE ASSESSMENT OF THE PARTY OF THE P

- It is herical value of it is locally incleared and of the mouse of the case.
- I depends on the same of T. and Y.
- The value of TC is calculated for one mole of a gas as 9TF using Acognitizes Law hazanting to Avapolities how At 9TP one male of a gas recuptes 22.414 drsf

900 51 mit. MATE

$$\frac{6 \times 10 \text{ mode}}{19 \times 2.414 \text{ deg}^4} = \frac{7 \times 0^9 \text{C} \times 2736 \text{C}}{8 \times 7} = 9 \times 1.0 \text{ min}$$

 $R = 62.4 \, \text{dm}^3 \, \text{mm} \, \text{of Ha mol}^4 \, \text{K}^4$

 $R = 62.4 \, dm^3 \, torr \, mot^4 \, K^4$ or

 $R = 62400 \text{ cm}^3 \text{ too mol}^4 \text{ K}^4$

P=1 atm =101325 Nm⁻⁴

T=273.16 K

n=1.0 mole

 $V=22.414 \text{ dm}^0 = 0.022414 \text{ m}^0$

$$R = \frac{PV}{nT} = \frac{101325 \times 0.022414}{1 \times 273.16}$$

1×273 16

 $R = 8.3143 \text{ Nm mol}^{-1} K^{-1}$ or $R = 8.3143 \text{ J mol}^{-1} K^{-1}$

Since 1 cal. = 4 18 J

$$R = \frac{8.3143}{4.18} \operatorname{cal} K^{-1} \operatorname{mol}^{-1}$$

R= 1.987 cal mot K1

Thus unit of 'R' are thus expressed in terms of " energy mol " R " !

- The value of 'R' in SI unit is $8.3143~\mathrm{J}$ mol* K^d , it shows that if one mole of an ideal gas is Present at 273 K and 1 atmospheric pressure and its temperature is increased by 1K, then it will absorb 8.3143 J of energy
- The value of 'B' in non-Si unit is $0.0821~dm^2$ atm mol⁴ $\%^4$. It shows that if one mole of an ideal gas in present at 273 K and 1 atmospheric pressure and its temperature is increased by IK, then it will absorb 0.0921 dm³ atm of energy. (dm³ atm is the unit of energy,

Example 8

A cample of nitrogen gas is anciosed in a sessel of volume 380 cort at 120°C pressure of 101828 Nm⁻¹. This gas is transferred to a 10 dm² flesh and cooled to 27°C. Calculate the pressure in Nm⁻¹ exerted by the gas at 27°C. (Guyanusia Board, 2008, Mulan Board, 2012)

$$V_s = 386 \text{ cm}^2 = 638 \text{ dm}^3$$
 $V_s = 10 \text{ dm}^3$

$$V_1 = 300 \text{ cm}^2 = 0.30 \text{ dm}^2$$
 $V_2 = 300 \text{ K}$
 $T_1 = 320\% C + 273 \text{ K} = 393 \text{ K} T_3 = 27\% C + 273 \text{ K} = 300 \text{ K}$

فقو أنحمده ود وبطعي

$$\frac{P_{\underline{z}} \nu_{\underline{z}}}{T} = \frac{P_{\underline{z}} V_{\underline{z}}}{T_{\underline{z}}}$$

$$\mathbf{Gr} \cdot \mathbf{P}_2 = \frac{\mathbf{P}_1 \mathbf{V}_2}{T_1} \times \frac{T_2}{\mathbf{V}_2}$$

Calculate the density of CH, (g) at θ^*C and 1 atmospheric pressure. What will happen is the density θ (a) transpositors in increased to 27^*C , (b) the pressure increased in 2 density θ of θ θ .

Off Calculate the density of mothems of STP (Labore Souri, 2010) Off Calculate the density of mathems at O'C and 760 mm Hg pressure. (D.G. Khan Soord, 2011)

$$T = 0^{\circ}C + 273 \text{ K} = 273 \text{ K}$$

Mol. manu of CH₄ M= 16 g mol⁻¹

Density is given by the formula:

$$d = \frac{1 \times 16}{0.0821 \times 273}$$

(d) Density at 27°C

Density is given by the formula

$$d = \frac{1 \times 16}{9.0921 \times 300}$$

of Density at 2 atmospheric pro-

Dennity is given by the formula

$$\mathbf{d} = \frac{\mathbf{PM}}{\mathbf{RT}}$$

$$d = \frac{2 \times 16}{0.0821 \times 273}$$

Example 5

Calculate the mass of 1 dm² of NH₂ gas at 30°C and 1000 mm Hg pressure, compidering that NH₂ to behaving theatly.

(Multin Board, 2012: Labore Board, 2014)

Solation:

0

$$P = 1000 \text{ rmm Hg} = \frac{1000}{760} \text{aim} = 1.316 \text{ aim}$$

$$V=1\,dm^2$$

$$T = 30^{9}C + 273 K = 303 K$$

Molecular Mass of NH₄ =M= 17 g mol ¹

$$R = 0.0821 \text{ dm}^3 \text{ atm K}^3 \text{ mol}^{-1}$$

Mass of the gas is given by

$$PV = \frac{m}{M}RT$$

0

0

College Charactery Ford-I

111

roles Q.P. (a):

What is Avogadro's loss of gases?

WOOADRO STAR

li states

t states

Equal volumes of all the passe
for of molecules (or moles)

So, number of moles of a gas is directly proportional to its volume.

- It shows that same number of moles of all gases occupies same volume at same temperature & pressure. The volume occupied by grie male of my gas at \$77 is called motor eclusive and is equal to 22,414 dec.
- One mole of any gas contains constant number of molecules. This number is called Associates's marriage. Its value is 6.02×10^{44}

$$\begin{array}{lll} \mbox{1 mole of Q_{2}} & = 32 \ \mbox{g Q_{0}} & = 22.414 \ \mbox{dm}^{4} \ \mbox{at STP} = 6.02 \times 10^{16} \ \mbox{molecules of Q_{c}} \\ \mbox{1 mole of H_{0}} & = 2.016 \ \mbox{g H_{0}} & = 22.414 \ \mbox{dm}^{4} \ \mbox{at STP} = 6.02 \times 10^{16} \ \mbox{molecules of H_{c}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} & = 22.414 \ \mbox{dm}^{4} \ \mbox{at STP} = 6.02 \times 10^{16} \ \mbox{molecules of H_{c}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g H_{0}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} \\ \mbox{2.6} & = 2.016 \ \mbox{g H_{0}} & = 2.016 \ \mbox{g$$

80.

I mole of O_{2} and H_{2} accupies some solume at STP, although O_{2} is 16 times heavier than H_{1}

It is because molecules in gases are widely separated from one another and have left evoply spaces. The distance between two molecules is approximately 300 times the dismest of the molecule. Therefore, size and masses of gas molecules do not affect their volume Hence, siqual moles of all gases occupy same volume at STP.

- One dust of every gas at STP will have molecules = $\frac{6.02 \times 10^{22}}{22.414}$ = 2.66×10^{61} molecules If temperature and pressure are changed equally for all gases, each gas will have $s^{ad} = 2.66 \times 10^{60}$ molecules.
- * One dm* of H $_2$ at STP watghs 0.0999 g. (since $\frac{2.016}{22.414} = 0.0999$ g) and one dm* of G_2 if

STP weighs 1.4384 g (since $\frac{32}{22.414}$ = 1.4384c) but they contain equal number molecules (i.e. 2.68×10^{61}).

Color Charleton Part I

11.

DALLON'S LAW OF PARTIAL PRESSERI

Debon's law states,

The total presence exerted by a minture of mon-reacting the partial presences of all the genes present to the mixture.

Udomanni Espression

Let partial pressures of different pases in a mixture are P_1,P_2,P_3 , then according to Delich's law, total pressure of this mixture is given by

$$P_i = P_i + P_a + P_a +$$

Parent Pressons The presence of each goe in a substant of games is collect percial pres

County and Explanation

Consider four cylinders of same volume each.

Three games H_0 , CH_4 and O_2 are enclosed appearably in three cylinders at the same imperature.

Let Pressure of $H_{\rm g}$ is 400 torr, pressure of $CH_{\rm g}$ is 500 torr and pressure of $O_{\rm g}$ is 100 lon. Let the three games are transferred to the fourth cylinder at the same temperature, then according to Dalton a Law total pressure of the mixture of gases will be

$$P_{\text{init}} = P_{\text{He}} + P_{\text{Obs}} + P_{\text{o}_{\text{c}}}$$

 $P_{\text{min}} = 400 + 500 + 100 = 1000 \text{ tory}$

- Since there are no attractive among the molecules of these guess in a mixture, therefore, svery gas exerts its own individual pressure. Hence, total pressure is the sum of individual ture of these gases.
- Due to independent motion of molecules, general gas equation can be applied to each gas Attendely. Thus

$$P_{H_2} V = n_{H_2} RT \text{ or } P_{H_2} = n_{H_2} \frac{RT}{V}$$

$$P_{OH_1} V = n_{OH_2} RT \text{ or } P_{OH_2} = n_{OH_2} \frac{RT}{V}$$

$$P_{O_2} V = n_{O_3} RT \text{ or } P_{O_3} = n_{O_3} \frac{RT}{V}$$

 $k_i T$ and V' are constants for all goses in a mixture

Therefore
$$P_{H_1} \propto n_{H_2}$$

$$P_{CH_4} = \pi_{CH_4}$$

College Characters: Protel

Hence, pressure of each gas is directly proportional to its number of motes.

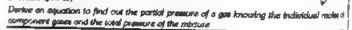
The total pressure of misture of pages will be

$$\begin{split} P_{t} &= P_{B_{d_{1}}} + P_{CB_{d_{1}}} + P_{D_{1}} \\ P_{t} &= n_{H_{0}} \cdot \frac{RT}{V} + n_{DG_{d_{1}}} \cdot \frac{RT}{V} + n_{D_{d_{1}}} \cdot \frac{KT}{V} \\ P_{t} &= (n_{H_{0}} + n_{DG_{d_{1}}} + n_{D_{0}}) \cdot \frac{RT}{V} \\ P_{t} &= n_{t} \cdot \frac{RT}{V} \end{split}$$

Where $n_i = n_{\mu_{ij}} + n_{\mathcal{O}_{X_i}} + n_{\mathcal{O}_{Z}} = \text{total number of all the gases}$

or
$$P_n V = n_n RT$$

This equation shows that the total pressure of the modure of games is directly proportion to the total number of moles of the gases,



I alculation of Portial Prosener of a few.

In a mixture of gases, purital pressure of any gas can be calculated if mass or moles of the gas total pressure and total number of moles of the gases are fundam.

Consider two games A and B forming a nebsure of games. Let estal preseure of the missure is P_1 and number of moles n_{ϵ} white. Partiel pressure of gas A is $P_{\rm A}$ and number of moles $n_{\rm A}$ and Partiel pressure of gee B is Pa and number of moles no

Then set can write

$$\begin{array}{ll} P_{a} V = n_{b} RT & & (1) \\ P_{A} V = n_{b} RT & & (2) \\ P_{A} V \Rightarrow n_{b} RT & & (3) \end{array}$$

Divide eq. (2) by (1)

MOLE PRACTION X

It is the number of holes of a substance divided by total number of holes of all the substances present in the mixture It is denote by X

see Ch#9, Solution#)
for some datails



Charleton Forts

$$\begin{array}{c} P_{A}V = P_{A}RT \\ P_{1}V = P_{1}RT \\ \hline P_{A} = P_{A} \\ P_{A} = P_{A} \\ \hline P_{$$

where, X_4 and X_6 are the mole fractions of gas A and B respectively. Thus, partial preson of a gas is equal to its mole fraction multiplied by the total pressure.

These equations can be used to determine the partial pressure of the green in a minute

Generally for I^{th} gas in a minimize of games, we can write $P_i = X_i \times P_i$

Example 6

There is a mixture of hydrogen, beliam and mathems compring a usual of volume 13 dm² at ST^*C and presume of 1 atmosphere. The mass of H_1 and H_2 are 0.2 and 0.12g respectively. Calculate the partial pressure in son H_2 of each gas in the interiors.

Solution:

Volume of the mixture of gases= V= 13 dm⁴

Temperature of the mixture $= T = 37^{\circ}C + 273 = 310 \text{ K}$ Pressure of the mixture $= P_{e} = 1 \text{ atm}$

Total no of moles = n_i = ?

R = 0.0821 dm³ atm K⁻¹ mol ⁻¹

Total number of moles is given by

$$P_tV = n_tRT$$

$$n_t = \frac{P_t V}{RT}$$

$$n_t = \frac{1 \times 13}{100} = 0.51$$

 $n_1 = \frac{1 \times 13}{0.0821 \times 310} = 0.51 \text{ moles}$ So, the lotal number of moles of H₂, He and CH₄ = 0.51 moles

Mass of $H_g = 0.8 \, \mathrm{g}$

Molar Mass of H_s = 2.016 g root⁻¹

0.8 g Number of moles of $H_a = \frac{0.8 \text{ g}}{2.016 \text{ g mol}^{-1}} = 0.396 \text{ moles}$

Mass of He = 0.12 g

Motar Mass of He = 4 g moi-1

Number of moles of He= $\frac{0.12g}{4g \text{ mol}^{-1}} \approx 0.03 \text{ moles}$

No of motes of $CH_a = Total males - (mole of <math>H_a + moles of He)$

= 0.51 - (0.396 + 0.03)

= 0.084 motes

Mole Fractions can be calculated as

$$x_{\rm H_2} = \frac{\rm No~of~oles~of~H_2}{\rm Total~Number~of~moles} = \frac{0.396}{0.51} = 0.776$$

$$x_{\text{His}} = \frac{\text{No. of moles of File}}{\text{Total Number of moles}} = \frac{0.03}{0.51} = 0.058$$

$$\kappa_{\rm OM} = \frac{\rm No. \ of \ moles \ of \ CH_{\odot}}{\rm Total \ Nmber \ of \ moles} = \frac{0.084}{0.51} = 0.164$$

Partial Pressures can be calculated as

$$P_{\mathsf{H}_{\boldsymbol{p}}} = \kappa_{\mathsf{H}_{\boldsymbol{p}}} P$$

 $= 0.776 \times 1.00 = 0.776 \text{ atm}$

$$= 0.776 \times 760 = 589.76 \text{ mm Hg}$$

 $P_{H_0} = x_{H_0} P$

 $= 0.058 \times 1.00 = 0.058 abm$

 $= 0.058 \times 760 = 44.08 \text{ mm Hg}$

Pou = xouP

= 0 164×1.00 = 0,164 atm

= 0.164 × 760 = 124.64 num Hu

Exercise Q10 (c):

Explain that the process of respiration obeys the Dalton's law partial pressure.

Applications of Dolton's law of Parint Pressure

Respiration process depends on the difference in partial pressures.



Callet Charles Parisi

+1+

Partial pressure of cooper in eir (159 torr) is greater than in the lungs (116 torr). Therefore, oxygen moves from air into the lungs. The pertial pressure of CO₂ in lungs is greater than in the air. Therefore, it moves out from the lungs into the air.

🕵 Breathing Ar High Aftitude

At sea level, the partial pressure of oxygen in air is 159 torr and breathing is easier. At At sea never, one passion pressure of oxygen is low it makes breaking difficult. That's why pilots use pressurized cable during flying.

3. Breathing In Deep Sea

Deep-sea divers cannot breathe with normal air in their tents. Instead, they use a misture of an inert gas and O₃ in which pertial pressure of exygen is adjusted around the sequired limits. Actually, in sea after 100 feet depth, the diver experiences 3 after pressure Thus, normal air cannot be breathed in depth of sea. Morsover, the pressure of N₂ increases in depth of sea and it diffuses in the blood.

4. Collection Of Green Green Water

Gases are usually collected over water. During this process vapours of $\mathcal{H}_{\varphi}O$ are mixed with the gas.

The total pressure ($P_{\rm annual}$) of this mixture ($H_{\rm g}O_{\rm mixture}+$ gae) will be

The partial pressure of vapous in gases is called aqueous tension.

P_{motet} = P_p + aqueous tension Thus,

P_e = P_{mote} - pqueous terulos

DRITESTON AND LETTINGON

Dillosum

0

4

The spontaneous intermiting of molecules of different games, due to collision, at a missionality and pressure is called gamesus diffusion.

Ny Inte office.

When different gases are mixed logether they wish to have some pressure everywhere. Thus, they go on mixing to form homogenous mixing until their partial pressures become equal everywhere.







er Characters, Partsi

1.15%

Emzple:

- The smell of rose or a scent spreads due to diffusion.
- The mixing of NO_g (a brown gas) and O_g (a colourless gas) is also due to $g_{\rm distance}$ diffusion as abown in the fig. The diffusion occurs by random motion and collision.

Huston

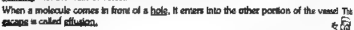
ft to the accept of you melecules one by one, without collisions through a hole of melecular size, into a region of loss pressure.



La porte de la

Why grown afficer?

The escape of gas molecules in not due to collision. It is due to their tendency to escape one by one. Gas molecules are habitual in <u>colliding</u> with the walls of vessel.



Exercise Q10 (d):

Ha

360 6	u do you differentiate between diffusion and effusion? Explain Graham's law of diffusion						
	Officence between Diffusion and Effusion of gases						
	16Bosien		[]Lps.on				
١	The spontaneous mixing of molecules of different gases is called diffusion.	1	It is the escape of gas molecules, one by one, without collision, through a hole of molecular size, into a region of low pressure.				
2	In this, gas molecules move from an axes of its higher concentration to an axes of its lower concentration.	2	It is due to the coiliding habit of mosecules with the wall of the container. During collision, when molecules come in front of a bole, they escape				
3	Gases diffuse by random motion and	3	Gases effuse without collision.				
1	Diffusion takes place in all directions.	4	Effusion takes place through a hole of molecular size				
5	Example: The smell of rose or a scent spreads due to diffusion	5	Example: Escape of gas molecules from puricipaed tyre.				



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Gerhan's Low of Diffusion

If was given by an English Scientist, Thomas Graham (1805 - 1869) The Graham s law states

At constant temperature and pressure, the rate of diffusion (or effusion) of a gas to inversely proportional to the equate root of its density or Molecular mass.

Mathematical Lagressian

 $r \propto \frac{1}{\sqrt{M}}$

Thus, lighter gases diffuse more rapidly than hearier gases.

Example:

NH₃ gas (molecular mass 17 g mol⁻¹) diffuses more rapidly than HCl gas (molecular mass = 36.5 g mol*)

Exploration

Consider two gases A & B.

For gas 'A' let its rate of diffusion in r_0 density 'd, , and molecular mass ' \mathcal{M}_1 ' For gas "B" let its rate of diffusion is r_s, density 'd_s, and molecular mass 'M_s' Then according to Graham's law

For gos 'A'

$$\eta \alpha \frac{1}{I}$$

$$\tau_1 \propto \frac{1}{\sqrt{d_1}} \quad \text{or} \qquad \quad \tau_1 = \frac{K}{\sqrt{d_1}} \quad \underline{\hspace{1cm}} \{1\}$$

For got B

$$r_2 \simeq \frac{1}{\sqrt{A_0}}$$

$$r_z \propto \frac{1}{\sqrt{dz}}$$
 or $r_z = \frac{K}{\sqrt{dz}}$ (2)

Constant K is same for all gases at same temperature and pressure.

Dividing (1) by (2), we get

$$\frac{r_1}{r_2} = \sqrt{\frac{d\epsilon}{d\epsilon}}$$

Strategy

$$\frac{n}{r_2} \neq \sqrt{\frac{M_2}{M_1}}$$

Where M_1 and M_2 are molecular masses of gas A and gas B respectively



College Chambris Park I

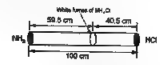
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O constraint of a gloring tracking street

Experiment

- Take a 100 cm. long glass tube, opened at both ends.
- Plug a cotton piece, soeked in NH₈ solution at one end and a cotton piece, soaked in HQ solution at other end.
- · The vapours of both these gases escape into the glass tube simultaneously.
- The vapours of both NH₂ & HCI are invisible. But when they meet with each other they
 produce white furner of NH₄CI.

Determine the distance travelled by NH₂ and HCl.



Calculations

Molecular mass of NH $_0$ = 17 g mol⁻¹ Molecular mass of HCl = 36.5 g real⁻¹ According to Graham's Law

$$\frac{r_{loo_1}}{r_{HO}} = \sqrt{\frac{M_{HO}}{M_{NH_0}}}$$

Thus

$$\frac{59.5}{40.5} = \sqrt{\frac{36.5}{17}}$$

1.46 - 1.46

Hence, Graham's law is vertiled

Francis 1

250 cm² of the emple of hydrogen affuses four times as rapidly as 250 cm² of si valueous gas. Calculate the inclar mass of enknown gas.

R remode Board, 2014. Sergodine Board, 2010

Charles Such

<u>Johnton:</u>

Rate of effusion undersown gas $= e_1 = 1$ Rate of effusion of H_2 gas $= r_{yz} = 4$ Motar mass of H_3 gas $= M_{yz} = 2$ g most $= M_2 = 2$ Motar mass of unknown gas $= M_2 = 2$

$$\frac{r_{H_2}}{r_k} = \sqrt{\frac{M_K}{M_{h_1}}}$$

Taking square on both sides

$$\begin{pmatrix} 4 \\ 1 \end{pmatrix}^2 = \left(\sqrt{\frac{M_X}{2}} \right)^2$$

$$\frac{M_X}{2} = \frac{16}{1}$$

$$M_X = 16 \times 2 = \boxed{32 \text{ g mol}^{-1}}$$

KINE IR. MOLLECULAR THEORY OF GASES.

To explain the physical behaviour of gases, a theory has been proposed known as kinetic Molecular Theory of gases.

1,11

- Kinstic Molecular Theory was proposed by Bernoull (11738).
- This theory was used by Clausius (2857) to derive the kinetic equation. He explained all the sess is the with this equation.
- Inwas further developed by Manuall, Boltzmann and van der want.
- Manuell gave law of distribution of velocities
- Boltzmann studied the distribution of energies among the see molecules.

The main pounts of this theory are

- All gases consist of very small particles called motorales. Gases like He, Ne, Ar have apprendic molecules.
- The inglecules of gases are widely separated form one another, therefore, these have large empty spaces between them.
- 3. The <u>Section</u> volume of the gas molecules is <u>negligible</u> as compared to the <u>total</u> volume of the gas.
- There are no attractive forces among the gas molecules. Therefore every gas molecule behaves independently
- The <u>average K.E.</u> of gas molecules is <u>directly</u> proportional to the <u>absolute temperature</u> i.e. Κ.Ε., α. Τ



- At the same temperature, molecules of every gas have game average kinetic energy
- Gas molecules are in generant modes motion. They change their direction only wh_{Ω} , they collide with one another or with the wells of container
- they collisions of gas molecules with one another and with the wells of container are perfectly elastic.
- 9 The presence of a gas is due to collisions of gas molecules with the walls of container.

 10 The motion given by force of gravity is negligible as compare to the motion given by

KINETIC CONTROL OF AN IDEAL TASK

- meta Egundana

R.J. Clausius derived the relationship for the pressure of an ideal gas known as kingle equation. It is given as

$$PV = \frac{1}{3} \, mN \, c^3$$

 $P = Pressure of gas, \ V = volume of gas, m = mass of one molecule of gas$

N= matrix of molecules of gas to the container, $e^{\lambda}=$ mean aquate velocity

Com Samo Section

سر جناه الله ايد س

If there are n_1 molecules of a gas with velocity ϵ_1, n_2 with velocity ϵ_2 and so on, then the a reposite velocity is given by

Movin square velocity =
$$e^2 = \frac{n_1c_1^2 + n_2c_2^2 + n_3c_3^2}{n_1 + n_2 + n_3 \dots}$$

Where
$$n_1 + n_2 + n_3$$
, $n_4 = N = total number of molecules.$

Under the given conditions, molecules of a get do not have some velocities. Actually event velocities are distributed among the molecules. Therefore, mean square velocity of e medacular a taken.

The expure road of matte square polectly to called root maps against sell by $a_{\rm min}$. The value of $c_{\rm min}$ has been calculated by using kinetic eq. it is given by



where. M= molecular mass of gas

This equation shows that higher the temperature careter the velocities of the case molecular

EXPLANATION OF GAS LAWS ON THE SASIS OF KIM HE MODEL COME THEORY. 1 Boule's Luc

Kinetic eq. for an xleal gas is

$$PV = \frac{1}{3} mNc^3$$
_____{{1}}

According to kinetic molecular theory of gases, the kinetic energy of ges molecules i.e. $\frac{1}{2}$ mNe³ is directly proportional to the absolute temperature, T

$$\frac{1}{2}$$
 mN $e^{\frac{1}{2}} = KT$ — $\{2\}$

Multiply and divide R.H.S. of eq. (1) by 2, we get

$$PV = \frac{2}{2} \times \frac{1}{3} mN_c^3$$

$$PV = \frac{2}{3} \times \left(\frac{1}{2} mN_c^3\right) \quad (3)$$

Pretting eq. (2) in eq. (3)

$$PV = \frac{2}{3}KT$$

If T = constant

Then PU- 1/

This is Boyle's low it shows that at constant temperatu * gap is impressly proportional to the pressure expried on it.

2, Darles & Law

Kinetic eq. for an Ideal gas is

$$PV = \frac{1}{n} m N_c^2$$



Colonia Charles Park

According to invote molecular theory of games, the average kinetic energy of g_{ij} metacodes i.e., $\frac{1}{2} \operatorname{molecular} h$ is directly propositional to the absolute temperature, "I"

$$\frac{1}{2}mNc^{3} \propto T$$

$$\frac{1}{2}mNc^{3} = KT_{---}(2)$$

Multiply and childs R.H.S. of eq. (1) 2, we get

$$PV = \frac{2}{2} \times \frac{1}{3} \text{ add } e^2$$

$$PV = \frac{2}{3} \times \left(\frac{1}{2} \text{ min } e^2\right) \underline{\hspace{1cm}} d 3$$

Putting eq. (2) in eq. (3

$$PV = \frac{2}{3}KT$$

Resettinging, we get

$$V = \frac{100}{30} \times 1$$

EP = constant then

This is Charles's less it shows that at constant presume, the volume of a given mast of one is discrete proportional to the absolute temperature.

CONSTRUCTOR CO.

Consider equal volume of two gases under the same conditions of temperature or maure.

Let - For gas 1 mass is m_1 , velocity c_1 and number of molecules N_1 ,

and Por gas 2, mass is m_{t_0} velocity c_0 and number of molecules N_0

Rinetic equations for the two genes will be

$$PV = \frac{1}{3} m_1 N_1 c_2^{N_2}$$

$$PV = \frac{1}{3} m_F N_E c_E^2$$

Since P & V are some for both quan



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Gener

Therefore

$$m_1 N_1 c^2 = m_2 N_2 c_2^2$$
 (1)

For both gapes, at the same temperature, the average K.E. per motocule is also some Thus $(KE_i)_t = KE_i t_k$

$$\frac{1}{2}m_1c_1^2 = \frac{1}{2}m_2c_2^2$$

.

Dividing eq (1) by (2), we get

$$m_1 N_1 c_2^2 \equiv m_7 N_2 c_2^2$$

Thus, equal volume of both gases under the same conditions of temperature an pressure, contain equal number of molecules, which is Avogadro's law.

Kinetic equations for the two gases will be

$$PV = \frac{1}{3} mN_c^2$$
 (1)

For 1 mole of gas $N=N_A$ and $N_A m=M=Moles ment of gas, therefore$



$$PV = \frac{1}{3}Mc^3$$

$$e^{x} = \sqrt{\frac{3PV}{M}}$$

$$PV = \frac{1}{3}Mc^{3}$$

$$c^{3} = \frac{3PV}{M}$$
Taking equate root
$$\sqrt{c^{3}} = \sqrt{\frac{3PV}{M}}$$

$$\sqrt{c^{3}} = \sqrt{\frac{3PV}{M}}$$

$$\sqrt{c^{3}} = \sqrt{\frac{3P}{M}} = \sqrt{\frac{3P}{d}}$$
since $\frac{M_{V}}{V} = d = density$

Root mean square velocity is actually the rate of diffusion of gas Therefore, all constant pressure

is shown that at constant temperature and pressure, the rate of diffusion (or effected) of gas is inversely proportional to the square root of its density, which is the Graham's law.

FINE DUENT RPRETATION OF TMPLRATOR

Consider the Minetic eq.

$$PV = \frac{1}{3}mNc^3$$
 (1)

m = mass of one motionale of gas. N. = number of motionales

V = volume

For one malecule the K.E. due to translational motion is given se

$$E_i = \frac{1}{2}mc^2$$
____(2)

E₀ = everage translational K.E. of motacides

Chalen: Pari

Muldaply and divide R.H.S. of eq. (1) by 2 we get

$$PV = \frac{2}{2} \times \frac{1}{3} \, mN_C^2$$

$$PV = \frac{2}{3} N \times \left(\frac{1}{2} m \zeta^2\right)$$
 (3)

Pat eq (2) in (3)

Let N=N_A = Avogadro's Number

$$PV = \frac{2}{3} \times N_A E_b \quad _ - \dots , 4)$$

According to general gas equation for 1 mole of a got

Compare (4) and (6)

$$\frac{2}{3} \times N_A \, \underline{P}_A = RT$$

or
$$E_k = \frac{3R}{2N_A} \times T$$

Thus there is a direct relationship between purefational K.E. and absolute temperature.

The temperature of a gas is directly proportional to the evenge bandational K.E. of its

Thus, a change in temperature changes the motion of motionies of a game

- The process of heat transfer from hot body to cold body is due to collisions of molecules with each other. During collisions, hat malecules transfer their $K.\Sigma.$ to colder molecules until the average translational $K.\Sigma.$ of the two bodies become equal. Thus, the T of two bodies had been separated by the first process of the process bodies becomes equal
- When T = 0 then E_K = 0 i.e motion of molecules stope, which is not possible. This importance is called absolute zero. It can never be achieved. The lowest temperature statinged to fair to 10 8 K.
- In game, and houlds, temperature is the measure of greature handstone. K.E. but in molids it is the measure of only ubrational K.E.



EIDDLEACHON OF GASES.

Corneral Principle of Liquelastion

The conventions of gases into liquids require high pressure and low temperature

- High P brings the molecules close to each other
- Low temperature decreases the K.E. of molecules. Thus, attractive forces are developed. among them and gas is converted into liquid

Copical Lemperature

The highest temperature at Which a substance can exist as a liquid is called critical

It a denoted by To

Carso of Pressent

re required to fiquidy the gas of the critical temperouses it calls The minimum pro

It is denoted by $P_{\mathcal{C}}$

Control Volume

The volume occupied by one male of got at $\hat{\tau}_c$ and $\hat{\mathbf{F}}_a$ is called Critical Volume It is denoted by Vo.

Example.

e For CO₂ T₀ = 31 1°C $P_{\rm ff}$ = 72.9 atm, $V_{\rm d}$ = 95.65 cm² mol *1

Importance of critical temperature

Pos every gas, there is certain temperature above which a gas cannot be liquefied. The temperature is called critical temperature

Examples, T_c for O_0 is 154.4 K, -...18.75°C). Similarly, T_c for CO_2 is 31.1°C. So, their gamm can be isquested only below their T_c . Hence, these must be cooled below that T_c and then recovered to T_c . and then pressure is applied to liquidy them.

Effect of Pelartsobility on artifical temperature

- Non-polar games have low polarizability and have a <u>very low</u> T_C e.g. T_C of Ar is 150.9 k. 1 122 26 °C1
- Paint several have high potentiability and comparatively high T_c 4.9, T_c of NH, is 405.6 $^{\rm K}$ (132 44°C)
- Thus, polar gases are easily converted into liquids

Salistiane 1	4 ritised temperature is that the beautiful for course				
Water Impaurs, H ₂ O	(41 6 37(.44°C)	2120			
Ammonia, NH ₃	405.6 (32.44 %)	103			
Prison-12, CCI ₂ F ₁	384 56°C	39 6			
Carbon dioxide, CO ₂	364,3 (1 42 ⁸ G)	79.0			
Окупеть От	54.4 8.75 C)	49,7			
Argen, Ar	50.9 (~122.26 °C)	41			
Nitrogen, Na	26. • 4° 96 °C)	33.5			

defluds for Liqueboction of Coses

Various methods are used for the inqualaction of gases. These methods are generally based upon uou's Thomson effect

both thomsauffeet

When a compressed gos is allowed to expand auddenly. It produces seeding. This is talked Joule - Thomson affect

Seasons in a compressed gas, moticules are very close to each other and have attractive force. When a gas is expanded auditinity moticules move away from each other. This process requires greeny, which is obtained from the gas itself itemer it is cooked.

Ind's Method

- * It's based upon Joule Thomson effect
- and liquefled air by this process.
- The compressed air (about at 200 aim) is passed brough a water-cooled pipe where the heat of compression is removed
- This compressed air is then passed through a spiral labe houling a jot at the end. When the gas comes out of jet into low pressure area. I aim i, it suddenly depends and is cooled due to soule. Thomson effect
- The cooled air moves up, cools the incoming gas of the
- del, and then again enters into the compression gamp, to be compressed again
- By repeating compression and expansion again and again, air a liquidisci
- All gases except H_a and He can be liquelled by this process



Engelie Q 13 (a)

35.

SON THE STEEL STREET, GASTA

help of a graph

While

The pin middle dove that other gree from each processor to called a "Real Circ" or a "Non-liked Gar

the properties of a term

The effect of temperature and pressure on behaviour of grees can be studied in ferror of pV = nRT

or PV = compressibility Sector

Samerally day

Thus value of componsibility factor is "1", for an ideal gas at all temperature and pressure.

That, for an ideal gas, when a graph is plotted between PU on Y-tota and P on X-tota a state of july to obtained parallel to commerce and

3. Caroni et Regions

Effect of Press

- ure, the curve starts somewhat along the ideal line. However, at very high P value of $\frac{PV}{nRT}$ has $\frac{1}{n}$ increased from the expected value and Fig.show deviation from ideal behaviour
- For H_b deviation starts over at low p



Office and Compared to the second of the control of

, For N_p PV first decreases below ideal line and then PV

CO₂ also shows unusual behaviou

Thes, it shows that deviations depend upon the patter of gas files of Temperature:

Al high temperature, the graph of these real gases come diser to the ideal line. Thus, the gases become kieal.

Conditions for Ideal and Non-Ideal behaviour of gones

- Gases are non-ideal at high pressure and law temperature.
- Generate ideal at way pressure and high temperature.

It obeys ges tows under all conditions of temperature and pressures. These ene no attractive or amount particles in regulgible accomplished forces accomplished forces among the ges particles particles in regiligible as compared to the joint volume of the ges. There are the attractive or among the ges perticles particles in regiligible as compared to the joint volume of the ges. The an ideal ges per ticles in the particles in the particles in the particles in the particles of the particles in the particles in the particles of the particles in the particles in the particles of the particles in the particles of the particles in the particles in the particles of the particles in the particles in the particles of the particles in the particles of the particles particles particles particles particles in the particles of the particles in the partic	
Contrato Q.19. (b). Do you think they	_
- Xul Mink that	

Do you think that some of the postulates of kirmic molecular theory of game are facilit? Point 44 here much facility? may your Derintons from (deptit)

Van der Wass (1873, pointed out that two ideal assumptions in the kinetic molecular temporarible for these deviations.

The actual volume of the gas particles is negligible as compared to the total volume of gas.

(ii) These are no attractive forces among the gas particles.

At high pressure and low temperature, gas molecules nome closer to each other and At high pressure and tow temperature, gas morecular come closer to each other and develop attractive forces among them. Moreover, at high pressure volume of a gas does he remain negligible So, the behaviour of real gases is not according to kinetic molecular heavy of gases and PV=nRT. Thus gases show deviations from ideal behaviour at be

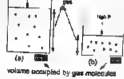
Exercise Q.14 (a)

Derive van der Waal's equation for real gases

can der Want's Equation for Real Gases.

van der Waat made volume and pressure correction to remove the defects of ideal pa mode! and gave an equation of state for real gases Volence Correction

Van der Waal pointed out that gas molecules have definite volume. Although volume of gas nave naminize volume. Although volume of gas molecules is very small as compare to vessel but it is not negligible. Thus, when pressure is increased on gas molecules, they oppose it So, if V is the volume of gas and 'b' is the volume of gas and 'b' is the volume of gas molecules per mole (exchaled volume), then the volume available for compression will be (V_max ~ b) and not V



Thus V_{bes} is the volume <u>available</u> to gas

Where constant b is the characteristic of each ges, its value is

"b" is the affective volume or excluded volume it is the volume occupied by 90 inducates in highly compressed state but not in liquid state.

Proposition Contract of

A molecule in the interior of gas has no unbalanced force it is because it is attracted equally from all sides. However, a molecule just striking the wall of the vessel is more attracted award. Therefore, it will not strike the wall with full force and hence observed pressure 'P' will be less than the ideal pressure 'P',' by an arround P



Consider two types of molecules 'A and 'B'. Let molecule of type A strikes the wall than it will be attracted inward by molecules of type 'B'. Therefore, P. depends upon the concentrations of A type and B type molecules. Le. P'α C_A C_B __(3

.

Let 'n' is number of moles of A an B present in a total volume "V" then concentrations of A and B will be given by

$$C_A = \frac{n}{V}$$
 and $C_b = \frac{n}{V}$ 14,

Put eq (4 In eq (3)

$$P' \propto \frac{n^2}{U^2}$$

$$P' = \frac{a \, n^4}{v^3}$$

For 1 mole of a gas

$$P' = \frac{a}{V^2}$$
 ___(5,

Where a = co-efficient of attraction or attraction per unit volumePor gases with strong intermolecular forces value of 'a' is high. Put sq (5) in eq (2)

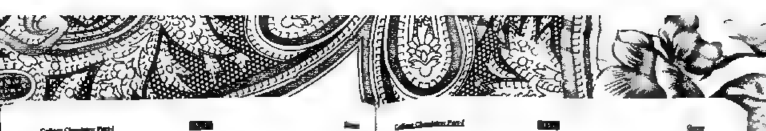
$$P_{i} = P + \frac{a}{V^{i}}$$
 _____ (6)

Thus, ideal gas equation for 1 mole will become

$$\left(P + \frac{a}{\sqrt{1}} \int (V - b) = RT$$

This equation is collect van der Waal's equation of state. This equation is applicable to real $t_{\rm trans}$

$$F_{OV \setminus p^d}$$
 (bole of a gap
$$\left(P + \frac{an^k}{V^d}\right) (V - ab) = nRT$$



complex. For H_0 gas $b=0.0266~\mathrm{dm^2~mol^{-0}}$ it means 1 mole of H_2 gas (2.016 g) occupies 0.0266 $\mathrm{dm^0}$ at clasest approach in gaseous state. For H_0 gas "of is the least (0.245) due to its non-polar character and small size.

15, 10

.

Units of b: m⁶ mot 1

Units of a:

$$\mathbf{n} = \frac{\mathbf{p}^2 \mathbf{v}^2}{\mathbf{n}^2} = \frac{\mathbf{N} \cdot \mathbf{m}^{-2} \left(\mathbf{m}^3 \right)^2}{\mathbf{mol}^2} = \frac{\mathbf{N} \cdot \mathbf{m}^{-2} \cdot \mathbf{m}^3}{\mathbf{mol}^2} = \mathbf{N} \cdot \mathbf{m}^4 \cdot \mathbf{mol}^2$$

Non-SI units

Units of b. dm⁶ mol * ⁶

Units of a

$$B = \frac{a_1}{a_2} + \frac{a_1 a_2}{a_2} = a_1 a_2 + \frac{a_2}{a_2} = a_2 a_2 + \frac{a_2}$$

The value of 'a' and 'b' can be determined by noting $\mathcal P$. V and T under two different

	restriction and their solu	activity to the major makes the same of
Lee	a diable to the first	South a reliable
hysioge.	0.443	U,07266
Onygo	+340	0.03 \$
Netropell	1 390	0.0391
Carbon decreids	3.590	0.0428
Caracia	6.170	0.0371
Bulgton districts	6 178	0.0564
Chlorist	6.439	0.0552

ed at 300 K. Ha column in 250 and, Calculate the Qua mote of ersed by the p

(0 When the gas in Ideal

(25) When the year to new-lebed

 $_{0} = 2.233 \, \mathrm{cm} \, \mathrm{dm^{2} \, mol^{-1}}, \, h = 0.0428 \, \mathrm{dm^{2} \, mol^{-1}}.$

Compains Score, 2012; Communic Scare, 2012; Labore Score, 2014)

$$n=1$$
 mole

 $R = 0.0821 \text{ dm}^4 \text{ atm } \text{K}^{-1} \text{ mol}^{-1}$.

According to general gas eq.

$$P = \frac{1 \times 0.0821 \times 300}{0.25}$$

in the goe to behaving as non

$$n = 1$$
 mole

 $R=0.0821~\text{dm}^3~\text{atm}~\text{K}^{-1}~\text{mole}^{-1},$

$$V = 0.25\,\text{dm}^4$$

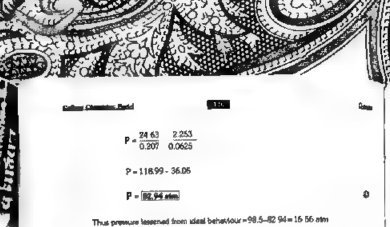
According to van der Waal's equation

$$\left(P + \frac{a_1 n^2}{V^2}\right)(V - nb) = nRT$$

or
$$\left(P + \frac{nn^2}{nt}\right) = \frac{nRT}{M_{A} \cdot nh}$$

or
$$P = \frac{nRT}{6(c-r)^3} = \frac{an^3}{c^3}$$

$$P = \frac{1 \times 0.0821 \times 300}{0.25 - 1(0.0428)} - \frac{2.253 \times 1^{1}}{(0.25)^{2}}$$



PLASMA 51.341

A unimpre of neutral particles, positive tons and negative electrons is collect Planne

consistence of Physics

- On heating a solid, it is converted into liquid. On further heating the liquid is converted into supposes. Thus, the phase of matter changes from solid to liquid and then liquid to vapount.
- Plant II vapous are further healed some of their lose electrons and positive ions are formed. Hence a resture of neutral particles, positive ions and negative electrons are produced. This is called Plantsis.
- The instanton is produced by high temperature or by radiations
- About 99% of the unsurers is made up of Plasma.
- If It is present everywhere in sun and start
- The sun is a 3.5 million km ball of plasma. It is heated by nuclear fusion
- It is the most abundant form of matter. It is the staff of stars
- it is present in everything from our to quarte (quark is the smallest particle of universe)
- Majority of the matter in inter-stellar space is plasms
- All the sharing main are plasma
- 8 On each, it is very limited, it is found in lightening bolts, finance, aucross and fiscretised lights, neon signs etc.
- When an electric outrant is passed through near gas, it produces both plasma and light

Callett Charleton Parts

137

Summer to the Cold Physical

- It consists of a significant number of charge particles. Thus it respond to both electric and magnatic fields
- 2 Motion of particles in the plasma produces fields and electric current within plasma density it refers to the density of charged particles.
- 3. Plasme has a complex set of interactions it is a unique factinating and complex state of matter
- 4. If is macroscopically neutral. Although it sontains sons and electrons but their number is

Sound and Archand Physics

- It is produced by using electrical charges on a gas e.g. in mean signs
- Placena at sow T is head to matritatin it is bacause outside a vaccium low T placena sacts rapidly with any molecule. Thus, it is both useful and hard to user.

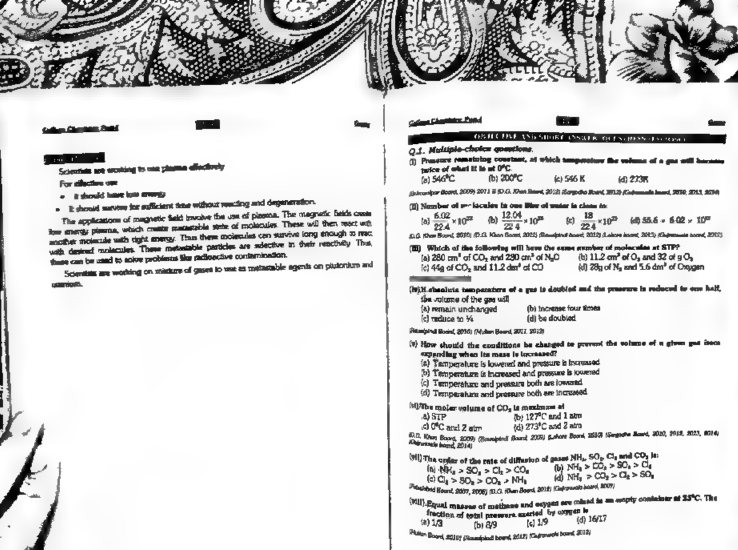
Natural Pleamo

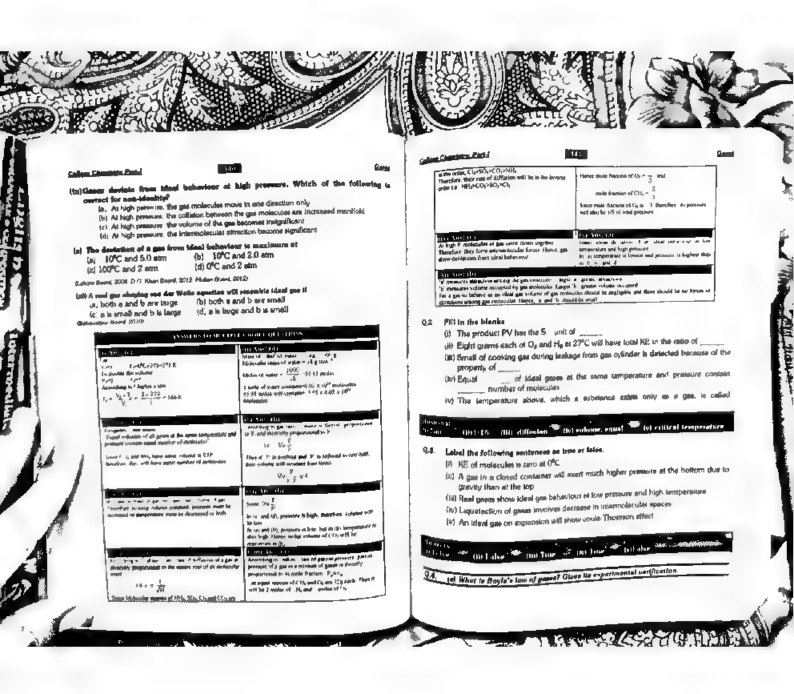
- · If only exists at high temperature or low temperature uncourse. If does not break down or react rapidly
 - These are extremely hot (over 20,000°C intrimum):
- They have so much energy, that they can vaportee any material

Indications.

Since plasma can respond to both electric and magnetic fields, it can have many uses

- 1. A fluorescent bulb is different from regular light bulbs. It consists of a long tube filled with gas. When startifely is passed through gas, it charges up the gas. The charging and exciting of gas creates glowing plasma inside the bulb
- Neon signs are glass tube filled with gas. When electricity is passed strough the tube. It thanges the gas and creates glowing plasma inside the tube. The colour of plasma depends upon the gas used.
- They are used for plasma processing of semiconductors, stardization of some medical Products, large, users, deemand costed time. high power microwave sources and Pulsat power switches.
- They are helpful for generation of electricity from fusion pollution control and removal of Reservous chemicals. Thus, it helps to clean up the environment
- 5 Plaume light up our offices, homes, it helps in working of computers, electronic equipment
- 6. It diffuse users and particle accelerators.
- i can be used to pesseurize lood.
- is in men to ways consists uniquel took







Carlos Charles Pard

Salved on Page 99

(ii) What are inotherms? What hoppens to the positions of leatherms when they are pleated at high temperature for a particular gos?

Solved on Page 100

(a) The do see yet a straight line when pressures exerted on a yet are plotted as inverse of volume? The straight line sharper its position in the graph by verying temperature dualify it

Solved on Page Ivi

(d) there will now emphase that the value of the acceptant if in (PV=10) in Heyle's less depend upon

(i) Temperature of the gas

According to Boyle's law PV=K at constant temperature. When temperature is increased volume of a given mass of gas is increased. Therefore, the product PV terms ly will also increase

(II) Quantity of the per-

According to Boyle's use PV=K at constant temperature. When quantity of gas it increased volume of gas is increased at consent temperature. Therefore, the product PV and hence K will also increase.

(a) When in the Chertae's four? Which scale of important is used to verify the V/T=0 (presents and number of moles are constant)?

Salved on Page 100

(b) A sample of certical memorates gas occupies 180 mL at 28°C, it is then asoled if constant pressure until II conceptes 100 mL. What is the new temperature?

 $V_1 = 150 \text{ mL}$

 $V_{\rm p}=100~{\rm mL}$

T1 = 25°C+273 = 298 K T1 = 3

According to Charles a law

 $\frac{V_1}{T} + \frac{V_2}{T_2}$

 $T_2 = V_2 \times \frac{T}{V_1} = 100 \times \frac{298}{160} = 196.7 \text{ K}$

(a) Do you think that the volume of any quantity of a gas become zero at -272°C to a new agents the less of accommission of mase? How do you deduce the idea of absolute sero free this information?

No volume of a gas cannot be zero at - 273°C since mass of gas will be destroyed in a gas of loss of consequences. genet the less of conservation of more which states

More sen metihar to arrested nor descriped

Com Chapter Park

141

Since temperature - 273°C is unattainable and it is the lowest semperature. Therefore it a plant as Absolute acco of Kelvin scale. Thus, absolute zero is defined as

The hypothetical temperature at which the softence of all goods has

(a) What to Kalein easily of temperature? Plot a graph for one make of an a real gas to prove that a gas becomes liquid, audior than $-273.16^{\circ}C_{\circ}$

Salved on Page 104

fij Throw some light on the factor 1/272 in Chiefae's law

According to quantitative statement of Cheden's law.

Al constant pressure, the volume of a given mass of an ideal gas increases or decrea by 1/273 of the original volume at 0°C for every 1°C rise or fall to temperature respectively

1. Thus, If we have 273 dm² of a gas at 0°C, then in $\frac{1}{273}$ part will be $\frac{1}{273} \times 273$ dm² = 1 dm²

to for every 1°C rise or fall in temperature, volume will increase or decrease by 1 dm²

f. Also, If we have 540 dm² of a gain at 0°C, then its $\frac{1}{272}$ part will be $\frac{1}{273} \times 546$ dm² = $2 \, \mathrm{dm²}$

to, for every 1°C rise or fell in temperature, volume will increase or decrease by 2 dm².

in both cases if temperature is decreased to = 278°C, the volume of the gas will become are Thus, = 273°C is taken as zero of katrin scale and is called Absolute zero. Hence, this fator 1/273 has helped in the development of Keivin Scale.

4.7. (c) What to the general gas equation? Darlos it in verticus forma?

Solved on Page 106

If Can two determine the molecular mass of an unknown gas V we know the pres temperature and solume along with the mass of the gas.

lobed on Page 107

Now do you justify from general gas squation that increase in temperature or ferroace of pressure decreases the details of the gas?

broad on Page 108

in the do see feet comfortable in supressing the donalities of guess in the units of plant. This is rether then glam

in the man molecular are unitally separated from each other, and small amount of the same and in a larger volume. Therefore, values of densities will be much wheller if express



Then in began and dist² in spall because depreciable emolati di gio sell la recession de and at the arts which at timelists will be grown the country of Chip of STP to 1 States and the special states and the states of the states of the special states of the states of the special states of the states of the special states of the states of the states of the special states of the states of the special states of the states of t

full. Dense the units for gameral gas constant if in general gas equation

m de de an

Thus
$$H = \frac{2^k \pi^{-k} \cdot m^k}{mod \cdot K} = 26m \mod 4K^{-k} = 3 \mod 4K^{-k}$$

n crange is segmented to says

to fill agreem. The units of R in spinus of always are $R=J \cos i ^{-1}R^{-1}$

in CGS system the amongs is expressed in sinist of any and

Transform using at R will be R = any and 190.1

However expension value of it will be delicant than St units

Q.D. (4) What is Assignative a law of ground

busyd in Figur 1.1

the the year stands when I made of M3 and I made of titls at GPL and I acce pro- house <u>designate</u> or symmetry of problems, if not judge

a sensor of more a service or some benefitive a some benefit of the sound of the services.

المقتلمين أند بيدودسون ويستههمون بيوسا جهاب الوس والة ليرس والأ لازد بلهوي الإستوب التسددا delife a 340th generation

to demandia alian it was not the west to raise of their as little will begin assure when application retractions are uncorrected and their to be being transfer alian place of light space of the

in games plantation trebusers have enclosure in departments 300 terms its midwooder tipe The reporter compared by the middle class flow and deposed often the respirators and an example. The measures The volume of one only despects upon the enclosure one or reason amount of the surples of second or property of the surples of second or second or of the surples of second or other or of the surples of t malicular although CFL reducials a Bassas teasure than He stitutes are

2.16 (at Daltum a law of partial promises to each silvaged by these partial and partial forms assume themselves Expirals it the publish days to

Deliver's and its on ideal gas law which assertes that there are no attractive terson-serving tio gas materiales. Planare away gan material masses statementalis.

On manifest different general if they have tagginglish adminishment for much retires these energy gas molecule mixture and particles of the property and transmission of the property of the property and transmission of the property of the

However if given develop strong forms for each other on interig, then their motivation of red for independent in their motion. Therefore presence asserted by each see self-for different their deposited. Hence great with deeping attractive transe do not vivry without a tea-

ld Desire up equation to find mix the partial prosours of a gas law value of comparisons green and the total pressure of the attenue Solved on Page 114

[9] Explois that the general of superstan aboys the Daine's law parties per Salant on Page 116

th New do som differentiate halance diffusion and affectour Exploie Graham s has at

listed on Page 118

9.18 (n) Game about the delay belonging at his temperature and high property Property this side the high of a parish

Stated of Proge 134

(b) The prior stands these arms of the procedure of bosons andorrow theory of grace are being that on the procedure. Subject and proper 1 41

ure, but SO2 and Cl2 are non-tiled (c) Philipping and Johan ere ideal at room inspesigned floard, 2007, 2010: Labore Board, 2019 How do you explain it?

 $H_{\rm g}$ and He have very low liquefaction temperatures. ($H_0 = -252.87^{\circ}C$ He = $-258.93^{\circ}C$ While SO₈ and O₆ have sufficiently high liquelaction temperatures, which are close to room imperature (SO₂ = 10.2°C Cl₂ = -34.6°C)

Due to very low liquefaction berspersture of Ha and He, their molecules have negligible attractions for each other at room temperature. Hence, these gases behave ideally at oom

While Equalization temperature of SO_{I} and Cl_{I} are close to room temperature, the of room temperature, their molecules have appreciable forces of attraction. Hence 50, and Cla are non-idual at room temperature.

Q.16 (a) Doriso may due Wool's equation for roof genera.

Salved on Page 132

(b) What is the physical algorithment of Vander Wall's constants a and b give their units (Molton Board, 2023)

In ven der Wash, equation 'a' is called as the co-difficient of attraction or attraction pt unit volume. It indicates the strength of Intermolecular forces in gases. Strenge to intermolecular forces, higher the value of 'a' and vice verse, its units are New ord 1.

In Van der Waaf's eq. % is called at the effective volume (excluded volume) compared by goe molecules in highly compressed sale but not in liquid state. It depends upon the set of gas molecules. Greater the size of gas molecules greater will be the value of b and we an. Six units one dan³ mot¹

Q 15: Explose the following facts

(i) The plot of PV serves P is a straight line at constant two perature and with a first ber of males of an ideal gas, (D.O. Khan Bourd, 2012: Acad Konberte Bourd, 2012)

According to Boyle's Law

At constant T, the product of printure and volume of a fixed mass of gas is all ant to PV = K

Hence, it is graph is plotted between P on X - tota and PV on Y-totis, then a straight life peralled to X – axis is obtained, showing that PV (or K) is a constant quantity

(8) The straight line in (4) to parallel to x-code and goes away from the pressure of ligher presence

74.

For rate gases, when pressure is increased, the value of PV does not remain constant. As higher pressure, molecules of gases come closer together, develop focus of attraction, and flus show deviation from ideal behaviour. Hence, at high P, the straight line changes to the arrow services in the pressure axis. The nature of curve depends upon the nature

(ii) The nen der Waals constant 'h' of a gas is four stress the motor volume of yes.

The gas molecules are incompressible spheres. Thus when gas molecules are pecked together like spheres, there remains some empty spaces in between the spheres which are

Hence, the incompressible volume 'b' is actually more than the actual recitar volume of gas molecules. Actually, it is four times than their molar volume.

a.e. $b = 4 V_{\bullet \bullet}$

(of Prossure of NH, gas at given condition (say 30 atm pressure and room temperature) is less as calculated by Van der Waat's equation than that calculated by general gas (Lithers Boord, 2012)

NH_a is a polar gas. Its molecules have forces of attraction. Thus, it shows non-ideal behaviour Therefore, malecules of NH₃ are estracted inward while striking the wall of container. Thus, these exert less pressure than expected ideal pressure

Hence, real pressure of NHs as calculated by Van der Waal's equation is less than the

(s) Water supouse do not belove ideally at 273 K.

(D.G. Khan Board, 2007: Lahore Board, 2012: Gujrentusia Board, 2013, 2014)

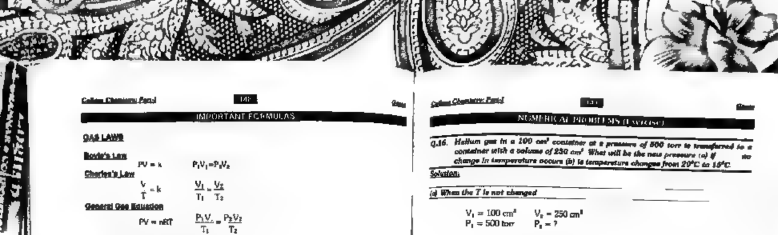
273 K is the freezing point of water. At this tersperature, vapours of water have **Porticiable forces of attraction between them. Hence water vepours behave non-ideally at 273 k.

(4) 60, In non-ideal or 273 K but behaves ideally at 387°C.

Ditulian Board, 2000: Labore Board, 2011, 2013)

At low temperature of 273 K, molecules 50, gas (10.2°C) have considerable the convergence of 275 K, necessary or an amount of the sound for each other and thus SO, gas behaves non-ideally.

When temperature is increased to 327 K, the forces of attractions among SO₂ molecules declarate and hence SO₈ gas shows ideal behaviour at 327°C



Bodr's Len

 $P_1V_1\!=\!P_1V_2$

Charles's Law

 $\frac{V_1}{T_1} = \frac{V_2}{T_2}$

General Gee Equation

 $\underline{P_1V_2} = P_2V_2$ PV = nRT

 T_1 $d = \frac{PM}{RT}$ $M=\frac{mRT}{pr_V}$

yan der WaaPe Equation

$$(V - nb) = nRT$$

intermediate

<u>Stronged rolls</u> Liver

To find the number of particles , atoms, lons, molecules or formula units number of particles = Given Mass × Na

or number of particles = moles $x N_A$

Mainr, Main may be Molecules mass or elemic regus or formula mass or ionic mass

Delten's Law of Bartini Pressure $P_{i} = P_{3} + P_{q} +$

 $P_{c}V \neq n_{c}RT$ or $P_aV = n_a RT$

 $P_i = \frac{n_2}{n_i} \times P_i$ or $P_a = \chi_a \times P_a$

 $F_p = \text{Total Pressure of minimum of gases}$ $P_p = \text{Partial Pressure of quests minimum}$ $\pi_p = \text{number of motion of a gas in minimum}$ $\pi_p = \text{Mote Praction of gas in minimum}$ $\pi_p = \text{Total number of motios of a minimum}$

er's Law of Diffusion or Effusion Of Gases

 $\frac{v_1}{v_2} = \sqrt{\frac{d_2}{d_1}}$

 $\frac{r_2}{r_2} = \sqrt{\frac{M_2}{M_L}}$

Charles Countly for an ideal Gas

$$PV = \frac{1}{3} \, m N_c^{\frac{1}{2}}$$

Q.16. Hallum gas in a 200 and consister at a pressure of 500 tors to transferred constainer with a volume of 230 and What will be the new pressure (a) if change in improving occurs (b) is temperature changes from 20°C to 15°C.

Solution

(a) When the T le not changed

 $\begin{array}{ll} V_1 = 100~\mathrm{cm}^0 & \quad V_0 = 250~\mathrm{cm}^0 \\ P_1 = 500~\mathrm{torr} & \quad P_0 = 7 \end{array} \label{eq:V0}$

According to Boyle's law

$$P_1 V_1 = P_2 \; V_2$$

$$P_2 = \frac{P_1 V_1}{V_2}$$

= 500 × 100

 $P_2 = \frac{500 \times 100}{250} = 200 \text{ fort}$

(b) When the T is changed

 $\begin{array}{lll} V_1 = 100 \ \text{cm}^3 & V_4 = 250 \ \text{cm}^4 \\ P_1 = 500 \ \text{term} & P_6 = ? \end{array}$

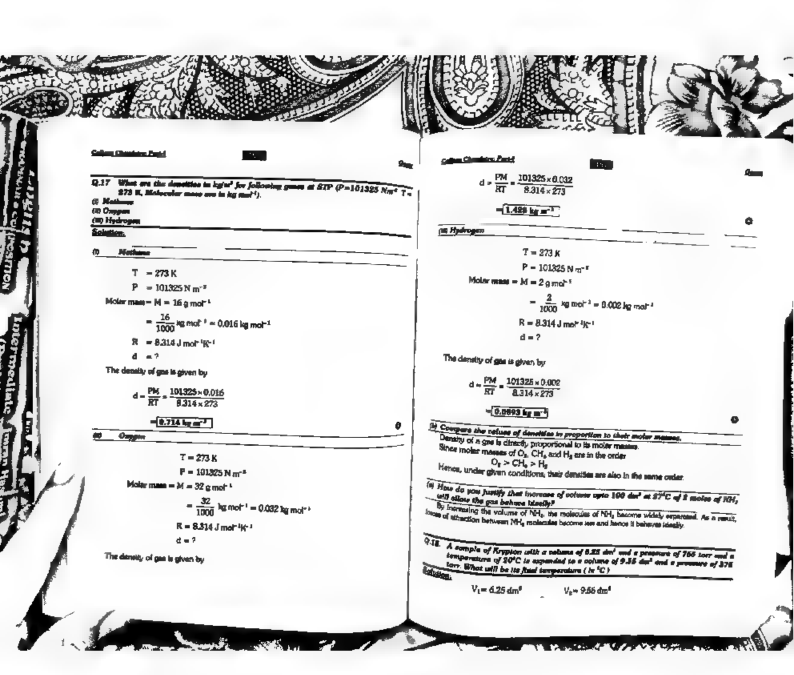
 $T_t = 20^{\circ}\text{C} + 273 \# 293 \text{ K} \qquad T_z = 15^{\circ}\text{C} + 273 \# 288 \text{ K}$

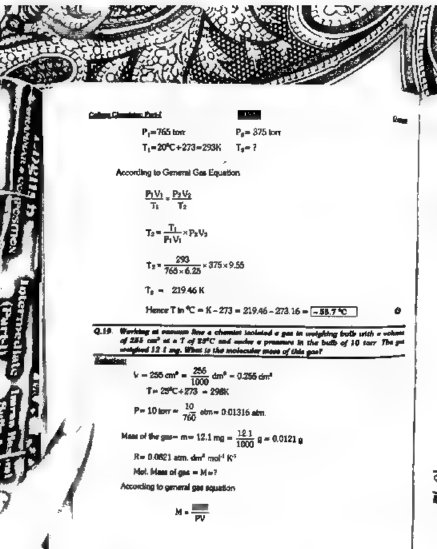
According to general gas equation

$$\frac{P_1 \, V_1}{T_1} = \frac{P_1 \, V_2}{T_1}$$

$$P_z = \frac{P_1 V_1}{T_1} * \frac{T_2}{V_1}$$

$$P_z = \frac{500 \times 100}{293} \times \frac{288}{250} = \boxed{196.6 \text{ term}}$$





161 0.01316 x 0.255 M = 68.25 g/mol Q.20. What presence is exerted by a mixture of 2 g of H, and 8 g of N, at 273 R in a 10 der? exemply est Mass of H_a = 2 g Males of $H_1 = n_{ext} = \frac{2}{2} = 1$ males Mass of $N_z = 8 g$ Moles of $N_a = n_{pq} = \frac{8}{28} = 0.286$ moles Total number of moles = $n_i = n_{ini} + n_{ini}$ = 1 + 0.286 = 1.286 moles Volume of mixture = V= 10 dm³ T= 273K 8= 0.0821 atm draft met 1 K1 Total pressure $= P_i = ?$ According to General Gas Equation $P_iV=\pi_i\,RT$ $p_c = \frac{n_b RT}{V}$ $p_s = \frac{1.286 \times 0.0821 \times 273}{}$ Q.21. (a) The relative denotities of two goess A and B are 1 1.5 Find out the solid B. which will diffuse in the same time to which 180 day of A will diffuse? Since relative densities of gases A and B are 1 1.5 Hence relative density of gas A = d_a = 1.5

And relative density of pas $B_{-} = d_0 \approx 1.5$ Volume of gas A diffused = 150 drs⁴



Internedial

Volume of gas B diffused = ?

According to Graham's Law of diffusion of gases

Since rate of diffusions are disuctly proportional to the volume of gases diffused,

$$\frac{\text{volume of gas B diffused}}{\text{volume of gas A diffused}} = \sqrt{\frac{d_A}{d_B}}$$

$$cr. \frac{\text{ we have of gas B diffused}}{150} = \sqrt{\frac{1}{1.5}}$$

values of gas B different
$$\pm \sqrt{\frac{1}{1.5}} \times 150 = \boxed{122.47 \text{ dim}^3}$$

hydrogen affects through a person plate at a rate of 800 cm 2 per estimate at 6° C in the rate of diffusion of expans through the same person plate of 6° C.

Rate of diffusion of hydrogen $= \epsilon_{ma} = 500 \text{ cm}^3/\text{min}$ Rate of diffusion of causers $= \epsilon_{ma} = ?$

Molecular Mass of $H_2 = M_{\rm tot} = 2~{\rm g~mol^{-3}}$ Molecular Mass of $Q_3 = M_{\rm tot} = 32~{\rm g~mol^{-3}}$

According to Graham's Law of diffusion of gases

$$\begin{array}{ccc} \tau_{OZ} & = & M_{H_{2}} \\ \tau_{HZ} & M_{OS} \\ \\ \text{or} & \frac{699}{500} = \sqrt{\frac{2}{32}} \\ \text{soc} & = \sqrt{\frac{2}{32}} \times 500 = \frac{125 \text{ cm}^{3} / \text{ ratin}}{2} \end{array}$$

(c) The rote of efficient of an induceum goe A through a plabele is found to be 0.275 than the rote of efficient of Hz through the same plabele. Calculate the motionics man.

 x_k Let Rate of allusion of unknown gas A = $x_k \approx 0.279$

Then Rate of effusion of unknown gas

Molecular mass of hydrogen = M_{HE} =2 g/mol

Molecular mass of unknown gas $A = M_A = 7$

According to Graham's law of effusion

$$\frac{nez}{r_A} = \sqrt{\frac{M_A}{M_{HZ}}}$$

or
$$\frac{M_A}{M_{H2}} = \left(\frac{r_{H2}}{r_A}\right)^2$$

or
$$\frac{M_A}{M_{H_2}} = \left(\frac{s_{H_2}}{r_A}\right)^2$$

 $M_A = \left(\frac{1}{0.279}\right)^2 n 2$

9.22. Calculate the number of molecules and the number each pro-

(Gujtararda Board, 2009)

(a) 20 cm² of CH, at 0°C and a pressure of 700 mm of Hg

 $V = 20 \text{ cm}^a = \frac{20}{1000} \text{ dm}^a = 0.02 \text{ dm}^a$ $T = 0^a \text{C} + 273 = 273 \text{ K}$

$$T = 0^{8}C + 273 = 973 K$$

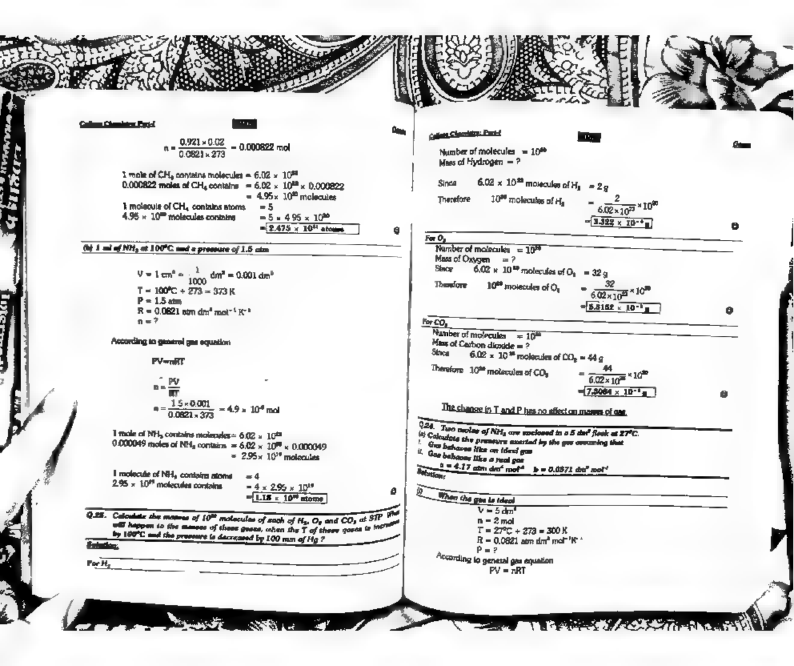
$$P = 700 \text{ mant of } \text{Fig} = \frac{700}{760} \text{ mins} = 0.921 \text{ mins}$$

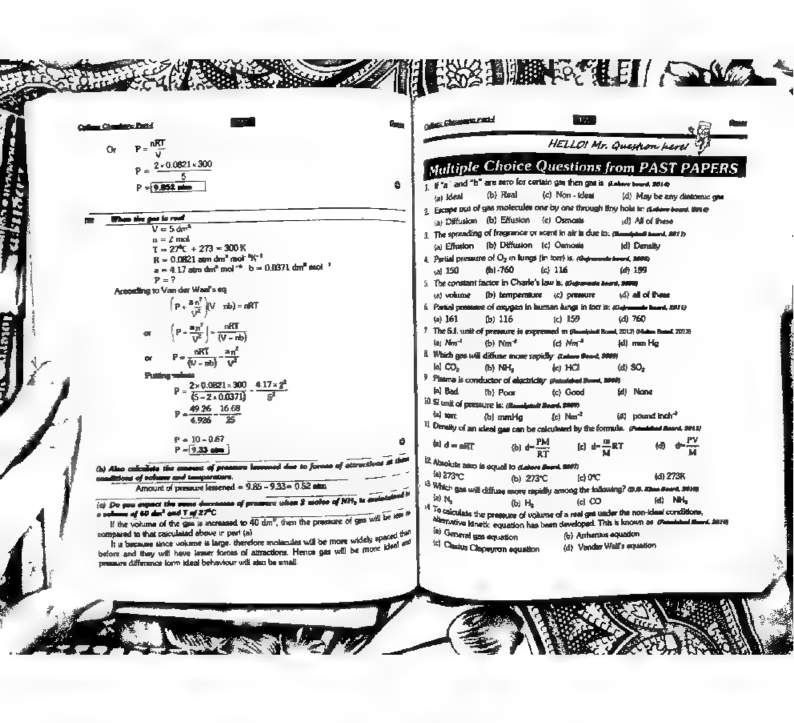
 $R = 0.0821 \text{ min dm}^3 \text{ mod}^{-1} \text{ K}^{-2}$

$$r = 1$$

According to general gas aquation

$$n=\frac{PV}{RT}$$









15. An ideal gas has volume 1 dm³ at 303 K. Keeping pressure constant, at which Keb_A, temperature its volume becomes 2 dm² (Sercotte Secret, 2011)

(a) 240

(ы) 308

[c] 330

160.

(d), 606

16 Which gas will diffuse more rapidly (Sugaran Band, 2014)

(a) CO₂

(b) NH₄

(c) HCI

(d) SO.

	Ansir	ers to 5	fultiple	Choire	Questra	lis from	Post Pr	DATE.	
Q#	Ane	Ο#	Аля	Q#	Ans	Q#	Ans	Q#	Ana
1	(a)	2	(b)	3	(b)	4	(c)	5	[d]
6	(b)	7	(6)	- 8	(b)	9	(c)	10	(c)
11	(b)	12	(b)	13	(b)	14	(d)	15	d
16	(b)								

Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK!

SHORT & LONG QUESTIONS FROM PAST PAPERS

PRESSURE, BOYLE'S LAW, CHARLES'S LAW

Short Overstone

(1 Define pressure. Gipe its units. (Lahora Bourd, 2009) Malian Bourd, 2009; D.O. Rhan Be Surgadus Bours, 2013) OR Compare the different units of pressure. (Surgadus Bours, 2007)

(2) Define Boyle's law. Give its expression. (Published Sound, 2009, 2012: Culturated Board, 2013) (3) The product of pressure and volume at constant temperature and number of motes t

constant, Why? (Sergodia Board, 2009) (4) Why the graph plotted between pressure and volume moves away from pressure at \$4

higher temperature. (Surgodia Board, 2011)

(5) Why do we get a straight line when pressure are plotted against inverse volume of a 95 (Behavelow Board, 2009: Gajamente Board, 2011: Revelopment Board, 2013)
 (6) Greater the temperature of the gas, closer the straight line of P versus 1N to the pressure on the straight line of P versus 1N to the pressure.

axis Justify It (Labora Board, 2007)

(7) State Charle's Low and write its mathematical form (Notice Board, 2018)
 (8) Volume of a gas is doubled when temperature is raised from O°C to 275°C. Wild (Submidum Board, 2008)

Define Boyle's law Give its experimental verification. (Behavely- Board, 2017)

48 OLUTE ZERO

- Sur Questions

 11. (Size the quantitative definition of Charles's law? (Panel guis)
- (2) What do you mean by absolute scro? (Fee dui Sourd, 2009: B and 2009: D.S. When Seved, 2019; QR What is absolute zero? What happens to real gases while approaching U? (Saryothe Board, 2012) Feliability Search, 2013)

(Bambbult Board, 2007 2009: Labors Board, 2014)

(Bambbult Board, 2007 2009: Labors Board, 2014)

SCALES OF THERMOMETRY

Mort Overthops

- Ment Streethers.

 What are different scales of thermometry? as What is thermometry? Name the scales and device used for thermometry. Wuten Board, 2011) OR How the derious scales of thermometry can be interconverted? (Laker Board, 1907)
- [2] Give two important scales of thermometry. How one these related? [Labora B Farelated Survey 2009]
- 3) 273°C is regarded as the lowest possible temperature, chiefly it. (General Board, 2010)
- (4) Valume of real gas connot be zero at any temperature. Why? (Surgodia Board, 2007)

(5) Convert 80°C to Fahrenhelt scale. (Februard Board, 1019) [6] Consert 40°F to Kelvin temperature Bahmalou Bassi, 2019)

7) Capuert 40°F to (a) Centigrade scale (b) Kelvin scale (a.o. 10mm Board, 2011)

(8) Containt -40°C to °F (Fahalahod Board, 2012)

[9] Convert 40°F temperature to keloin temperature (4 mobile Band, 2012)

ENERAL GAS EQUATION IDEAL GAS CONSTANT

Short Charitiens

I) What is general gas equation? Derive it in various forms, phases home, 2010

(i) Derive the unit of 'R' in general gas equation when the pressure is in atmosphere and Derive the unit of 'R' in general gas

(3) Coloniate the value of R to SI units OR Derive the units for gon constant R in general gas equation (alten pressure is in New and volume in M (Labors Board, 2005, BSIS) Guyamada Seed, 2011, 2013, 2014 Milliam Board, 2006, 2017; D.G. (Can Seed, 2009, Reselpted Board, 2009, Rule: Sept. Board, 2012, 2012)

(4) Derive general gas equation for one mole of a gas? (Outramoula Board, 2011)

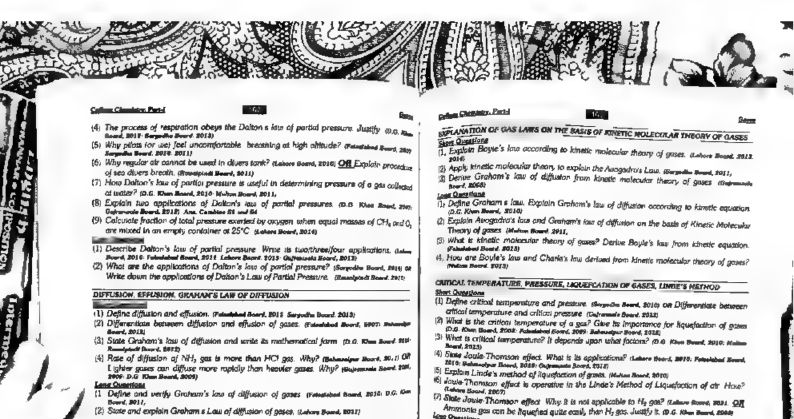
AVOGADBO'S LAW, DALTON'S LAW OF PARTIAL PRESSURE AND ITS APPLICATIONS

Observing Grant Land of gases? (Roundard Band, 2010: (Supposite Band, 2016: Labore Board, 1010).

What is Apogadro's law of gases? (Roundard Band, 2010: (Supposite Band, 2016).

Sand, 2021: Behavelar Beard, 2021) OR Define Dakon's low of partial pressure. Give Politican Control of C

(3) Probe that $P_1 = P_1 X_1$ (Labore Board, 2014)



KINETIC MOLECULAR THEORY OF GASES

ad. 2009)

1) List four postulates of kinetic molecular theory of gases. (Familian Bord, 2011, takes

(2) Give mathematic expression of Kinetic equation and root mean square valuable. (No

(1 What is Kinetic molecular theory of gases. Give its Postuicies, (Sequebe Borni, 2013)

| A) Why guess do not settle down in a vessel? (Gurennorie Board, 2008)
| 5] What are elastic collisions? Give an ecomple, Vestom Board, 2007)

(3) Ove mathematical expression for mean square velocity and root mean square velocity

Short Ounsidens

1000 Ownstiens

erc. 2012)

Less Observious
(1) Describe _under's method of liquefuntion of gases. (Labora Seast, 2013)

ther Countries

(b) What is compressibility factor? What is its value for an ideal gas? Multim items, 1007.

(c) What is compressibility factor? What is its value for an ideal gas? (Febulahad Boses, 1008).

[2] What are faulty points in kinetic molecular theory of good? (Fatalabad Bown, 2009; Lad

The point deed behaviour (manufacture) and the production of the production of the pressure and tout temperature make a gas non-ideal? Off Gases deviated this pressure and tout temperature make a gas non-ideal? Off Gases deviated the pressure and pressure and the pressure and t

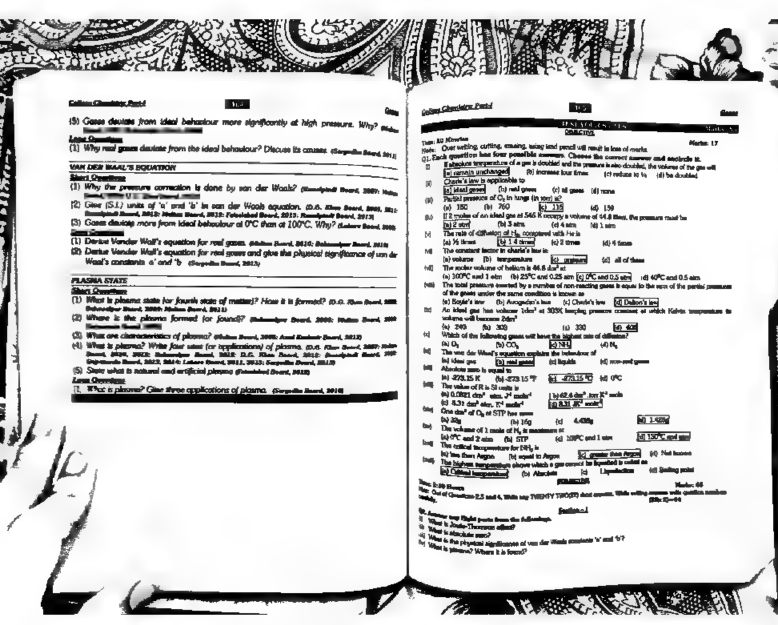
significantly at high pressure and low temperature, they wise Sound, 2007 1810: Guy

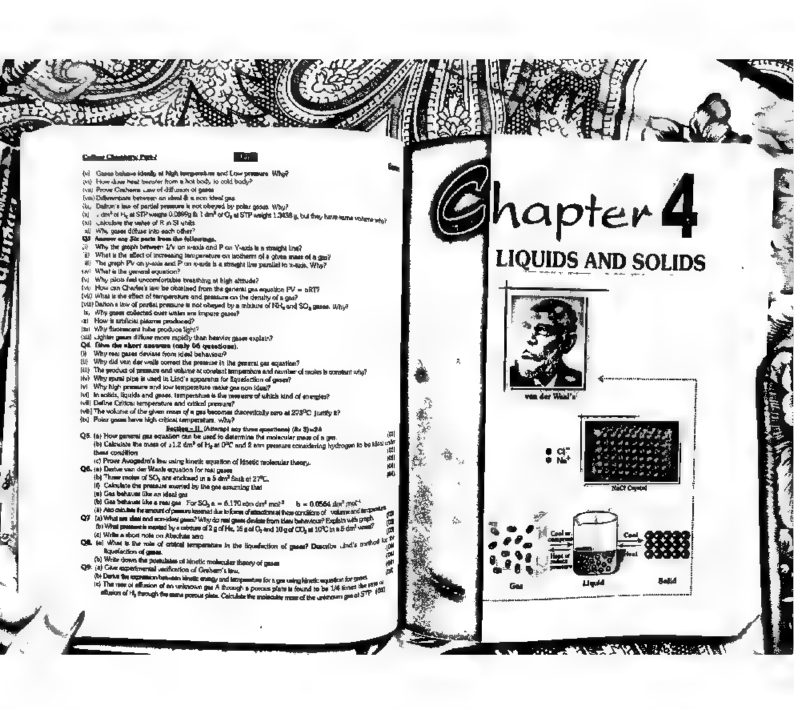
ad Board, 2013, 2019: Malein Board, 2014: Multan Board, 2913)

hand, 2009: Behaveler Board, 2011; Labors Board, 2015; OR Describe two causes of deviation From Indexty Companying Sourd, 2015; Lancer process, 2015; Lancer

DEAL AND NON-IDEAL GASES, CAUSES OF DEVIATION

(4) 2050; Pole







CONTENTS .

Chapter-4

- Liquids and Solids

INTERMOLECULAR FORCES

Dipole-dipole forces Dipole-induced dipole forces or Debye forces instantaneous dipole-induced dipole forces or Landon dispersion forces Factors effecting the London forces Hydrogen bonding
Properties and applications of compounds Containing hydrogen bonding

EVAPORATION

Vapour pressure Measurement of vapour pressure of a liquid Bolling point Bolling point and external pressure Energetics of phase change Energy changes and intermolecular attractions Change of state and dynamic equilibrium LIQUID CRYSTALS

Objective and Short Answer, Questions (exercise) INTRODUCTION

Types of solida CRYSTAL LATTICE CRYSTALS AND THEIR CLASSIFICATION Properties of crystalline solids CLASSIFICATION OF SOLIDS

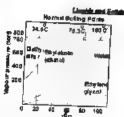
ionic solida Covalent solida

Moiscular solids Markillo edilda

DETERMINATION OF AVOGADRO'S

NUMBER (NA)
Objective and short answer, questions (exercise)
Post Papers MCOs and Short Questions

165



Remobble HON

Liquids have indefinite shape but definite volume

in liquids, intermoterular distance is greater than solids but less than gates and intermotecular forces are stronger than gases but weaker than solids.

INTERMOLLS DEAR AND INTRAMOSCOPER PORCES

Internal centur Forces

The fugues of distraction between separate mole monotoxilar forces

- These forces are called van der Waals forces.
- These forces are present among all types of atoms and molecules when they are close to each other
- The physical properties of substances e.g. melting point, builing point etc. depend upon the strength of intermolecular forces
- These forces have no selation with valence electrons.
- These forces are weaker than intramolecular forces.

Emples.

Dipole-Dipole forces; London dispersion forces etc.

The forces of entr Committee,

Chemical bonds i.e. conic bond, covalent bond, co-ordinate covalent bond.

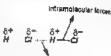
They have almost no relation with physical properties of substances.



Liculate and Solute

- These forces have concern with valence electrons
- These forces are stronger than intermolecular forces.
- Intramolecular forces are stronger than intermolecular forces.

Consider HCl. In this, a covalent bond is present between H and Ct. Both atoms complete their valence shell. Thus, they tend to always remain together. Hence, this linkage is very strong While, in intermolecular forces only a many altermotatic attention of the contract only a weak electrostatic attraction is present between Of of one molecule and H^{ab} of other molecule. So, this linkage is weak.



Types of a molecular Farces

Intermolecular forces are also called van der Waals forces, especially when the molecules are close to each other

There are many types of intermolecular forces

Following are the common types

- 1 Dipole dipole forces
- 2. Ion dipote forces
- 3. Dipole induced dipote forces
- 4. Instantaneous dipote unduced dipole forces or London forces

The electrostatic forces which are developed when a regative end of one polar molecule attracts the positive end of other molecule are called dipole-dipole forces.

by ample and Explain man

- Consider a bond between two different atoms e.g. H Cl. In HCl. Cl is more electronegative than H. It attracts shared μ . electrons more towards itself. Thus, CI gets partial negative charge (δ^*) , while H gets partial positive charge (δ^*) .
- Such a bond is called polar bond and the molecule is called a dipole.
- So, when molecules are close to each other they tend to line up and attract each other However, thermal energy of molecules does not permit perfect alignment
- These forces are approximately one percent as effective as a covalent bond

Factors offering Dipole Dipole faces

The strength of dipole-dipole forces depends upon

College Chambelon Pathil

17 [

. Ofference in electronegativity of the bonded stone

Greater the electronegativity difference, more polar is the bond, hence, stronger the dipoledipole forces and vice versa

Example: Dipole-dipole forces in

Distance between molecules.

arger the distance between molecules, weaker the dipole dipole forces and vice versa

Exemple: an gases, molecules are widely separated, therefore, these forces are very weak.

in liguids, molecules are close to each other therefore. these forces are stronger



Part of Dipole Dipole forces on Physical Properties

- The physical properties of substances e.g. melting point, boiling point etc. depend upon the strength of dipole-dipole forces.
- Generally, stronger the dipole-dipole forces greater the values of thermodynamic properties of a substance like melting points, boiling points, heat of vaportation and heat of sublimation etc.

Digrac Books - Constant - Constant I - Constant

The electrostatic forces of attraction between permanent dipole of one molecule and tableed dipole of another molecule are called dipole-induced dipole forces. Descripto

These are present in a mixture of polar and non polar molecules. In this mixture Positive and of polar motecule attracts mobile electrons of non-polar motecules.











the Burning applica-

Thus, dipute is induced in non-polar molecules. The force of offsetion between polaries. molecule and induced dipote is called dipote-induced dipote force or Debue force.

incontaneous Dipole-Induced Dipole Forces or London Dispersion Forces

The maintaining force of attraction between instantaneous dipole and induced dis-sited instantaneous dipole - induced dipole force or Landon dispersion force.

Non-polar He gas can be liquefied which shows that its molecules have forces of Meaching. To account for such things. Fritz London, a German physicist, explained these in 1930. These are called London dispersion forces.



These forces are more prominent in non-noter molecules e.e. E. N., noble once etc. in These forces are more promunes as passesses increases the dipoles. However, in negroods molecules, dipoles are not present under normal conditions.

- Consider non-polar He molecules. When two He molecules come closer to each other, their electrons repet each other and are pushed away. Therefore, a temporary dipole is produced in the molecules. Thus, electron density of molecule is no more symmetrical and it becomes a dipole. This is called justimismeous dipole
- The positive end of instantaneous dipole attracts electrons of another molecule. In this way, a dipole is induced in the nearby molecule, as well. This is collect induced discle.
- The force of attraction is then developed between instantaneous dipose and indeed dipole it is called instantaneous dipole-induced dipole force or London dispersion total
- These forces are developed only for few moments because electrons are making
- continuously.

 The temporary dipole is finished very soon. However, a new dipole will appear in some other direction and thus weak forces are again developed between molecul
- These forces are present in all types of molecules (polar and non-polar). However these are more prominent in non-polar molecules e.g. H_B, Cl_B, noble gases etc.
- London forces are weaker than dipole dipole forces.

Salanzahilda

ment of the extent to which the electron cloud our be dista sed is called Polarizability.

A specie (atom, ion, and molecule) is said to be polarized if temporary dipoles into occasion of electron cloud.

Lacturs allering Fundin Luccs

Size of Heatronic Cloud

Larger the use of atoms and molecules, discontion is easy and polarizability is in Hence London forces are stronger.

Caller Classician Part I

Married and South

1

Exemples:

Boiling points of Noble Gasse and Helogens Increases Down The Group in Periodic Table

Noble gases are monatomic gases. They do not form covatent bonds among themselves because their outermost shells are complete.

In Noble gases, electronic cloud size increases down the group in periodic table due to increase in atomic size. Thus, atoms are easily polarized down the group, and develop strong London forces

That's why boiling points of noble gases increases down the group.

Similarly boiling points of halogens (VI) A, also increases down the group.

Halogens have different physical states at room temperature.

Halogens are non polar diatomic molecules.

F₂ and Cl₂ are gases. Br₂ is liquid while l₂ is solid

It is because, electronic cloud size increases down the group in periodic table due to intrease in atomic size.

Thus, atoms are easily polarized down the group, and develop strong condon forces. Hence physical states of halogens changes down the grap and boiling point also increases, e.g. boiling point of F_a is $-88.1^{\circ}\mathrm{C}$ while boiling point of rodine is $+184.4^{\circ}\mathrm{C}$ it is because colorization. polarizability of codine is much greater than fluorine due to larger size.

¹⁷ Q 34.6

Smaller of Atoms in a Mali cule

Generally, greater the number of atoms, stronger the London forces and vice versa,

Greater number of atoms means greater polarizability and house stronger London forces

" Ac 485.7 35 Br ™ Kr 152.3 58.8 Se Se 184.4 107 L ¥ Sin 61.6

WIIA

188.1

VIII A

268.6

P No

-245.9

Share (C.H.) have lower boiling point (-88.6°C) than because, C.H., (68.7 °C).

Although both C₄H₆ and C₆H₁₄ are non-polar However, due to larger number of atoms in C₆H₆. It has stronger forces of attraction than C₆H₆.

It is because, target hexane molecule has more places for attraction and greater laneautes. polarizability than smaller whene molecule has more proces for assumed and higher boiling.

Point Bridge than smaller whene molecule 50, it has stronger forces and higher boiling. DONE Further, due to stronger forces, hexans is a liquid while ethans is a gas at room authouse in a



Physical state of hydrocarbons changes with increasing mulecular mass

Physical states of hydrocarbons changes from gas to liquid to solid with increasing Physical states of dynocentors and places for attraction and great molecular mains. It is because larger molecules have more places for attraction and great potanizability than smaller molecules. So they develop stronger forces, Hence, melling an boiling points also increases with increase in molecular masses.

Table 4.	2 Burbay p	aints and p	hysical states o	af same þy	drocarbons
Name 3	a, BP C (Laim).	Physical Matteal STP	Normal M	Majori Park D	all refer ad
Methane	16.0	QE3	Penlane	361	biopes
Ethons	-88.6	Ges	Hexans	68.7	Liquid
Propone	42 .	Ges	Deceme	174.1	Liquidi
Butane	-0.5	Gas	[sedecane	327	Solid

Hydrogen Banding

The electrostatic interaction between n a strong electronegative atom and path positionly charged hydrogen atom is called hydrogen banding

The strong electronegative elements are mostly N, O. F and rarely CI

Two spolecules involved in H-bonding may be some or different

J. cample & Laplanation

Hedrogen boading in H.O.

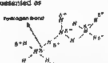
Consider H_eO. In H_eO, strong electronegative O attracts shared pair of electrons more towards itself. Thus O gets large 5 charge and H gets large & charge Thus, dipole-dipole interaction may be developed among water molecules. However losses of attraction among water molecules are stronger than simple dipole-dipole interaction

Oxiden has two lone pains of electrons. Moreover "H" creates a strong electric field due to its small size. Thus, the expension atom of one H₂O molecula limits to the H-atom of another H₂O molecula limits to the H-atom of another H₂O molecula limits to the H-atom of another H₂O molecula ione pair by a <u>co-ordinate covalent bond</u>. This bond formed is called <u>hydroden bond</u>

Hydrogen banding in HF and NH₄

Hydrogen bond between HF and NH_a molecules are represented as





16.1

Hydrogen bonding in occions and chloroform

Some elements other than N. O. and F. are also involved in H bonding, e.g. in also highly 5°. This H-atom can now tomb H. bend with strong electronegative atom of other molecule e.g. with paygen atom of actions as shown in the lig.



Properties of Dudrogen Hand

- Hydrogen bond is tonger than normal covalent bond.
- Hydrogen bond is stronger than dipole-dipole interaction but weaker than normal covalent bond. It is generally 20 times weaker than covalent bond.
- Hydrogen bond is a directional bond.
- Hydrogen bond results in the formation of long chains and network of molecules.

Properties and Application of Compounds Containing Hedrogen-Bonding

Strength of Acids

HF Is a Weaker Actd than HCl. HBr and HI

In HF, molecules are H-bonded in a zigzag manner Thus, H is entrapped between two F aloms as shown in the fig.

Thus, HP cannot easily donate its H1 rons easily, hence

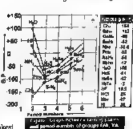
A increased manie Properties of Caralese Medilles

In covalent hydrides of Group IV A to VII A. H-bonding effects are very clear

Consider the graph for boiling points of covatent hydrides plotted against their period numbers.

Soling points of hydrides of Group IV A me lovest among all covalent hydrides.

is because elements of groups IVA are seasi electronegative, therefore, they have weakest intermotecular forces among all covalent hydrides, e.g. CH_a has towest boiling point because it is a very small motecular. Molecule and its has least polarizability





The boiling point of NH_a, HF, And H₂O are Highest in Their Respective Series.

It is because these hydrides have strong electronegativity elements N. F and O, with form hydrogen bonding among their own molecules. Thus, their bolling points are high

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H₀O has High B.P. Than HF, Although F is More E.N. Than Q.

It is because, F atom can make only one H-band per molecule due to presence of the hydrogen, while H_eO can form two H-bond per molecule because it has two hydrogen atoms and two lone pairs of electrons. Hence, due to presence of strong hydroger bonding in H₀O. its boiling point is greater than HF

Botting point of NH₁ is lower than HP and H₂O.

NH_a can form one H-bond per molecule. It is because N has only one lone partit electron. Also, its electronegativity is lower than O and F. Hence. Its boiling point is lower than HF and H₂O

H₂O is a liquid but H₂S and H₂Se are goes

in H₂O strong H-bonding is present which makes it a liquid. In H₂S and H₂Se was intermolecular forces are present. Thus, H₂S and H₃Se are gases at room temperature.

Boiling point of HBr is Higher Than HCI

It is due to bigger size of Br than Cl

Due to bigger size of Br, HBr has greater polarizability and stronger London tons among its molecules than HCI. Hence, boiling point of HBr is greater than HCI.

The hydrides of fourth period e.g. GeH., AsH., H.Sc and HBs show greater bolling points than those of third period due to larger size and greater polarizar littles.

Exercise Q5 (RI):

water and ethanol can mix easily and in all proportions.

salabilate

Both H₂O and Ethyl Alcohol (CaH₂OH) are Highly Muchle talth Each Other

Substances, which can form H-bond with each other are highly soluble into each other. Since both H₄O and ethyl alcohol can form hydrogen bonding with each other therefore, they are miscible with each other in all proportions.

However, larger alcohols are not soluble in water due to non-polar nature of bigger hydrocabon chain in them.

Similarly, small carboxylic acids (RCOOH) are also soluble in H₂O

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Hydrocarbons are macluble in testas

It is because they are non-polar and cannot develop H-bonding or other attractions with H.O. Hence, hydrocarbons are insoluble in water

Heaving Action

Soops and detergents are made up of long non-polar hydrocarbon tall (generall, allyl or being)] and a polar anion head. In water, the head is stabilized by making H-bond with H.O. while non-polar tail remains outside H_eO because it is not soluble in water

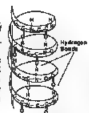
Thus, hydrogen bonding helps in cleansing action

J. O Fermina, a Biological Compounds and Land Materials.

H-banding is very important in living organisms.

Large Protein malecules in Itsing organisms are stabilised des to H-bonding.

Many fibrous proteins e.g. horn, nail, skin, feather, hair etc are composed of long chains of amino acids. These chains are collect around each other and form a spiral. This spiral is called help. Such hells may either be right handed or left handed in f@ht handed helix groups like NH and C=O are vertically adjacent to one another and they form H-bonds. These Nicingen bonds link one spins to the other



Note analysis hove shown that on the everyor there are 3.6 amino acids for each tun of the helps.

DNA (decorribonately); acid) occurs in cells.

It consists of two spiral chains which are collect about each other on a common sate than Louble hells. This is 18-20 A* in diameter. They are linked together by hydrogen bounding between thair eublinits.

E. Bombou in Paints and Doos

The adhesive nature of pertain paints and dues is also due in Hbonding with the surfaces.

Striderty, sticky action of Give and Honey is also due to H-



2, and Material

Food material such as carbohydrales e.g. glucose, fructose and acrose are also stabilized due to H-bonding. All these contains OH stap which produce H-bonding.



College Chambers: Part-I

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Armide and Toke

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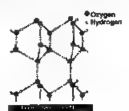
Both natural and artificial fibres have rigidity and tensile strength due to H-bonding.

H_eO has tetrahedral electronic structure

fee Florie on water. In liquid HaO, molecules form temporary H-bond with each other it is because due to movement of molecules, bonds are broken and reformed. Hence there is less regularity and less free space.

However when temperature of H_0O is lowered below $4^{\circ}C$, its molecules become gauge and form permanent H-bond. So, empty spaces are developed in-between the molecules and its volume increases. Ice occupies 9% more space than limits water. Thus, density of its becomes less than water. Hence, it floats over water

Similarly Setumes assecture of Ice and dismond: The <u>structure of ice</u> is just like that of a disposed because each storm of carbon in diamend is at the centre of <u>intrahedran</u> just like the <u>concert</u> of water molecule in <u>ice</u>. It has becaused structure with large empty spaces



Application of few density of too to cold elimates: Density of ice is iss than water. Therefore, in cold climate, when temperature fells below 4°C, cold water being lighter come to the <u>surface</u> and feezzes to ten. Thus, an <u>insulating says</u> of ice is formed above gain to the surface. They was a feeze to the property of the same of the property of the same of the Heavy The sayer of he prevents further heat loss from undermeath water. Thus, it protects acuatic life from cold.

Chemistry Parts PARTIFICATION

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Literalds and Solids

The spontaneous change of a liquid into aspoure is called evaporation athuous at all temperatures. it cor

According to kinetic molecular theory, molecules of liquid posses idnetic energy. All the Moderalises do not have some K.E. Some molecules have K.F. higher than the average webs.

When such type of molecules come to the surface of liquid, these overcome the resmolecular force and thus escape from the surface of liquid as vapours. This is called evaporation.

In an open container, at constant imperature evaporation continuous at the same rate until all the liquid is converted into vapour. Evaporation continuous at all temperatures

Le has Allecting the Rate of Leaguentain

The rate of evaporation of a liquid depends upon the following factors.

Strength of intermulecular Porces

Deferent liquids evaporate at different rates.

Stronger the Intermotecular force. Lower will be the rate of exaponation and pice waste e.g. at the same temperature, rate of evaporation of gasoline (petrol) is more than water ecause petro: has weaker London forces

Similarly rate of evaporation of water is lower than other. Because in water strong hydrogen bonding is present.

2. Temperature

increase in temperature increases the note of emporation and vice versa. It is because, increase in temperature increases the number of molecules having K.E. higher than the average value. Thus more molecules are escaped from the surface of liquid, Hence rate of evaporation increases

Surface Army

Larger the mericoe ones, higher is the rate of evaporation and vice versa, it is because, pare number of molecules escape from larger surface area. That is why expanded dothers are dried earlier than unexpanded dothers.

orities Q12: Explain the following with reason. (§ Evaporation causes cooling

Ecoporation Course Cooling

emperature is the measure of average kinetic energy (K.E.) of the particles of a Ribetance. Greater the K.E. higher is the temperature.

When particles having high KE escape from the surface of liquid, the average KE, of Then particles having high K.E. escape from the surface of industrial decreases. As the emporation particles decreases. So, temperature of the Equid decreases. As the emporation continuous, liquid absorbs most and most heat from the surroundings, for temperature of the surroundings, and the surroundings of the surroundings. harmone, temperature of the supporteding also decreases. Hance, evaporation causes tooks.



Color Chartery Part 1

180

Lincoln and Su

Emercine Q12: Explain the following with reason.

to. Dynamic equilibrium is established during evaporation of a liquid in a closed vessel

VAPOUR PRESSURE

The pressure exerted by the support of a liquid, in equilibrium with the liquid, α , an important to called support pressure of a liquid.

STATE OF THE PERSON NAMED IN

J

When a liquid at placed to a closed container, then, due to evaporation, molecules lesson the surface of liquid, and change into the vapour state. Some of these amount on colliding with other victours, tone a part of their K.E. and are

minural to the liquid state. This is condensation process.

instally the rate of evaporation is higher than the rate of condensation. After sometime, when the space above for legaci becomes seturated with apours, evaporation rate becomes equal to representen the and a dynamic emailingum is established as the system

Pogram Enterkomment of dynamic American propagation and include

Like gas, supposes of liquid also collide with the wells of container and event pitt et art angelifference in eatherd response presentate of the Septiet.

The expert presence to independent of the amount and surface were of the house.

The assignment is famor from larger surface area. However, solutioning molecules dis-tures imager area to subarri. These, condensation is also faster 50, number of molecules po-nets surface area remains some. Thus, vapour pressure is independent of amount and surface. ause of the based.

Fillowing factors affect the vapour pressure of a housed

belonge the interviolecular forces lower will be the sopour pressure of a liquid and vice versa.

Example I

At 35°C H₂O with strong intermolecular hydrogen bonding has low vapour pressure (43 mm of Hg) than ester (442 mm of Hg) in which so hydrogen bonding in

Falde 4-3 Majorian Arang Aphyland	C 41 20
Home of compound	Vagorial (ACE)
Exhyl other	580 442.2
Chlossiann Cadon Tarachloride	120
Carron 1 mrscmores	419
Kercury Glycosol	G 00015

Colonia Company Part of

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Балужа 2

soperizate has the high trapour pressure due to weak Leaden forces, while glycerol has the low due to strong hydrogen boarding.

increase in temperature incremes the papers pressure of a liquid and vice persa.

It is because at high temperature, R.E. of molecules pagests, therefore, evaporation increases, Hence, vector measure increases.

Example

are of H₂O increases from 4.579kerr to 9.209 and from O°C to 10°C, but it increases from 527.8 tour to 760 torr from 90°C to 100°C

The rate of chance of vaccur pressure with temperature agree by Chances—Canceyron equation.

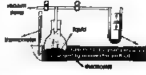
ru	to show
0	4.579
20	9 209
20	17.54
383	31.42
37	47.07
40	56.32
50	92.5L
60	149-4
30	233.7
40	355.1
90	527.E
100	700.0

TOTAL STREET

less trement of Vajour Presente of a Lapid.

nomenic Method

- This method is used to determine accurate vapour pressure of Squid.
- Appearable is not as shown in the fig.
- Place liquid is placed in the round bottom flock and is freem.
- Aix above the liquid is removed with a man часных разпр.
- Fixen liquid is remaited to release the
- Liquid is again frozen and air above it is This process is repeated at the air is removed.



- The fleak is then kept in a thermostet at a temperature at which vapour pear liquid is to be determined
- The supposes of the liquid pushes the speccary column in the extenometer. After formatimes height of mercury in both limbs of manameter becomes constant.

The vapour pressure of the liquid is given as

P =P + Ah

where P = pressure of the liquid, P_{--} = atomospheric pressure, and

4h = difference in height of Hg in manometer



Little and Sales

BOILING POINT

A TOP A

The temperature of which the popose pressure of a liquid becomes equal to the external pressure to called bottling point of a liquid. Explanation

The vapour pressure of a liquid socreases with increase in temperature. When a liquid is heated its vapour pressure goes on increasing until it becomes equal to the external pressure. At this temperature, liquid starts boiling. This is the boiling point of liquid.

The amount of heat required to reporter one mole of a figured at its b.p. is called Moles heat of reporteration.

At boiling point bubbles continuously come out of the boiling liquid. It is because, bubbles formed in the interior of liquid have higher internal P than atmospheric pressure. Thus, the bubble come out of the liquid and burst upon the surface.

Table, Bading Parals of some commute liquids							
Lagurds	BP O	Liquids 7346	407.701				
Apetic wold	118.50	Carbon tetrochloride	76.50				
Acetone	56.00	Elihamol	78.26				
Andine	184.4	Maphthalene	218,00				
Repaire	80 15	Phenot	181.60				
Carbon disulphide	46.30	Water	100.00				

Effect of intermolecular forces on boiling point

Consider the vapour pressure curves of water, ethyl alcohol, ethylene glycol and diethyl ether

- Graph shows, that all the liquids boil at a temperature, at which their vapour pressure becomes equal to 760 torr at sea level.
- At some temperature, curve of diethyl other starts a 200 test, while that of water at 4.8 test it shows that the to weak informolecular forces, other has higher vapour pressure than water

Thus, ether boile at lower temperature than water.
Hence weaker the intermolecular forces, lower will be the

 The curve shows, that vapour pressures for all liquids increases yet, ranch, closer to the houting point of itentia.

Empire - Vagnatu Liverstein alle Community of the Shared Scientific Calent Chambers: Port-d

Bading Point and External Pressure

The beffing point of a board increases with the increase in external pressure and vice pros

980

It is isocause, at higher external pressure, higher needs to absorb more heat to equalize its vapour pressure to external pressure hence boiling point is higher. While at fower external pressure liquid needs to absorb less heat, hence boiling point is lower.

e.g., at 760 mm of Hg water boils at 100°C, while

at 1489 mm of Hig water boils at 120°C

Exercise Q12. Explain the following with reason.

(all) The boiling point of water is different at Murree hills and at Mount Everes

Water balls at lawer temperature on Marree hills and Moura Everest.

On Momee hills, external atmospheric pressure is round about 700 mm of Hg, which is lower than normal. Hence, water bolls at a lower temperature of 98°C, At further higher althide, on Mount Everest, external atmospheric pressure is further discreased upto 323 mm of Hg, hence water bolls just at 69°C.

Applications of Vaccation of Boiling Councients Presser.

The variation of botting point of a liquid with pressure has many applications a.g., Pressure cooling, vacuum distillation at:

d) Parsogge Caroking

Policible: It is based on the fact that increase in exactnal pressure increases the boding point distinuit.

figuretance: At higher altitudes, abmospheric pressure is lower than normal. (1 am). Thus water boils at lower temperature e.g., it boils at 95°C at Murree hills (P=0,92 atm). Thus cooking takes more time.

The holling point of water can be reised in a pressure cooker if is a closed container in which vapours are not allowed to escape. Consequently they develop more pressure on the water. Flence, boiling point of water is increased. This halps in cooking the meet & registables quickly even at high altitudes.

English Q12; Explain the fellowing with resson.

My Nacyum distillation can be used to avail decomposition of a sansitive liquid.

2) Voc. and Distillation

he distinction corried out under reduced pressure is called second distillation.

The first is based on the fact that decrease to pressure decreases the boiling point of a

Importance: Some high boiling liquids carnot be apparated by distillation at normal pressure lie, I such, since they decompose before reaching their boiling points, e.g., glycerine boils at



Lincoln and Saltin

290°C but decomposes before reaching its boiling point. However when the extend pressure is reduced to 50 mm of Hg in a vacuum distillation apparatus, it boils at 210°C and therefore can be distilled without decomposition.

UNURGETICS OF PHASE CHANGE.

During physical or chemical changes, energy is evolved or absorbed.

Luffinge Change

The energy charge at constant P, in a physical or chemical process to called Enthaly,

It is denoted by ΔH It is expressed in kJ/mox

Three types of enthalpy changes are associated with physical changes

Modor hear of fusion

It is the amount of heat required to convert one mole of a solid into liquid at the ing point

It is denoted by A.H. It is expressed in lel/mot

Mida hem of equalization

Is to the amount of heat regulared to compart one main of a liquid into exposure at its boiling point.

it is denoted by A H., It is expressed in kel/mol

Moder heat of subfinement

It to the amount of heat required to convert one mole of a colld into expoure at its aublimetten neint.

It is denoted by ΔH_0 , it is expressed in kV/mal

France Changes and Intermolerular Attractions

Enthalpy change of a physical change a.g. repartization can determine the strength of

When a liquid is heated, its temperature goes on increasing until its boiling point is reached. At boiling point the temperature is stopped. Now the hear supplied is used to cover the liquid into vapours by breaking intermotecular forces,

The amount of heat required to convert one mole of liquid into vapoure at its boiling point is called motar heat of vaporization (A H₉).

Generally, Higher the value of A He, stronger the intempolecular forces and vice with Thus, AH_V is actually a measure of strength of intermolecular forces.

185

11 1 10

~ 40.6

4 21 7

+ 56

+ 24 6

+ 5.9

+ 10.00

+ 45.00

+ 27.00

+ 8.60

+ 15 .

+ 16.9

+ 30.1

Scheduler

H₂O

HQ

SO.

£,

O,

84

CH₄

C₂H₄

C,H,

 C_nH_{ns}

Sourcise Q12: Explain the following with reason.

(to) Heat of sublimation of a substance is greater than that of heat of vaporis

Best of Sublimation to Larger than of Heat of Vaportestion

Fleat of vaportzation is the amount of heat required to convert iquid into vapours, while, heat of sublimation is the amount of heat required to convert solid into vapours.

Since intermolecular forces are stronger in solids than liquids. therefore, it is difficult to vaporize a solid them a liquid Hence, heat of sublimation is larger than beat of vaporization.

Heat of Vaportzation	Footen	đ	Substances	Бę	Less	than	Heat	o,r
			r					

Heat of fusion is defined as the amount of heat required to convert a fixed amount of a solid into liquid at its m.p.

While, hear of vaporization is defined as the amount of heat upped to convert a fixed amount of a liquid into its vapours at its boiling point

To convert solid into liquid, intermolecular forces are not greatly broken, but to convert liquid into vapours, intermolecular

forces are largely broken. Therefore, vaporization requires more energy than fusion

Hence heat of fusion is less than heat of veportration

Starcios Q12: Explain the following with resson.

(kg) Heat of vaporisation (or subtimation) of todine is very high

ls has Higher Heat of Sublimation them its Family Members Subst Polerizability has stronger intermelection forces w with greater

Polarizability increases with increase in etomic size. Since la has considerably larger size Then its family members, therefore, it has greater polarizability, and hance along intermolecular forces. Due to this, it is difficult to vaporize 4. Hence it has higher heat of whitements. aublimation.

Polar Substances have Higher Values of $\Delta H_{\rm F}$, $\Delta H_{\rm S}$, $\Delta H_{\rm f}$

Polar molecules have stronger intermolecular forces, thus large energy is required to Charge their physical phase form solid to liquid or liquid to vapours er solid to vapours

Hence, polar substances have higher values of ΔH_0 , ΔH_0 , ΔH_0

⁶8. H₄O, SO₄, NH₃ etc are polar substances and have considerably higher value of A H₂.



Colon Classics, Park

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Charles and Sulph

Charge of State and Distantite Equilibrium

Sanctific and con-

The process in which both forward and reserve process can copy me sale, executable processes.

Dynamic republicant

The stage of reservible process, at which rate of forest d charge becomes equal to ρ_0 and attachment charge to called an equilibrium stage

Since both changes are occurring simultaneously at equal rate therefore this equilibrium is called dynamic equilibrium.

Example.

At O°C, solid ice exists in dynamic equilibrium with water

ice

ofc.

-

CIOUTO CRYSTALS

The intermediate turbid liquid phase of some solids between solid phase and disliquid phase, shouling some properties of liquids and some properties of solids is said liquid crustal.

Lift, aid Crystele were first <u>placovered</u> by an Austrian botanist. F. Reinitser, in 1988, he was studying an organic compound <u>cholesterul benzoate</u>. This compound becomes <u>milital</u> 145°C and becomes <u>clear at 179°C</u>. On cooling reverse process occurs. This turbid mily phase was called liquid crystal.

Profession .

When a solid is melted, it is converted to a clear liquid.

However, many crystalline solids pass through a turbid liquid phase before firsh conventing into clear liquid. This <u>turbid obese</u> is called <u>liquid crystal</u>.

A louid crystaline phase exists between two temperatures, a meiling temperature clearing importance

Malifest Incomment

Charles arrangement

Solid == Liquid Crystel == Clear Liquid

As professor English Crestalls

Their properties are intermediate between crystals and isotropic liquids

- I These have some degree of order like solids.
- 2. These have <u>fluidity</u> like liquids.

Colon Congress Parts!



- 3. These have properties such as <u>surface investor, process</u>; etc. like hands
- 4. These have obside properties of crystalling made
- 5. These are 3000000

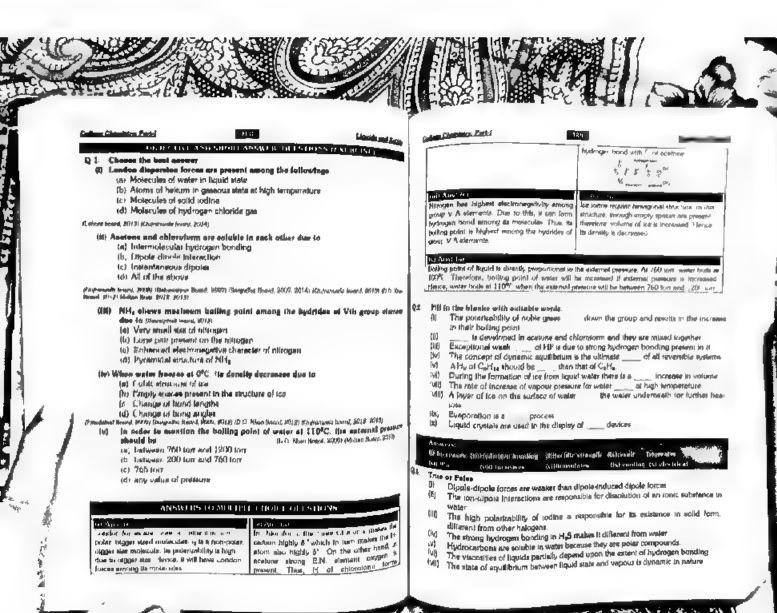
Types of Liquid Crystale:

On the basis of orderma of paracles, house creates can be divided into someons know a a

pplication

From 1988 to until about 30 years ago, liquid crystals were limited to laborationies. But now these have many applications

- I Equid crystals have excellent electrical and optical properties
- 2. Many organic compounds and biologica, tissues behave as liquid crystale
- 3. Liquid crystals gan diffract tight. When one of the visvetengths of light is reflected by crystals they appear coloured. With change in temperature, the distance between layers of molecules of liquid crystals also changes. Hence, colour of reflected light also changes. Thus, liquid crystals can be used as emperature-sensor.
- With change in temperature, liquid crustals change colour and give measure of temperature. Thus, these are used to prepare room thermometer, for a deligite temperature range.
- This can be used to find the point of potential follows in electrical circuits.
- 6. Liquid crystals are used in medical diamonia. These can be used to locate veins, affections and turnours it is because these parts are warrier than surrounding issues, so they can be detected by using temperature sensitive liquid crystals. This technique is called <u>Skim thermosphophy</u> it is used.
 - (I) to detect blockages in veins and arteries.
 - (iii) to detect the breast cancer at earlier stage is layer of temperature sensitive liquid crystal is pointed on the surface of breast. The warm lumour shows up in blue colour.
- 3. These are used in display of electrical devices <u>if CD = iterate region display</u>, such as digital watches, calculators and computes it is because temperature, pressure and electromagnetic fields can change the week bonding conditions to cayutais, hence give different appearances.
- In <u>chromatooraphy</u>, liquid crystals are <u>used to solvents</u>
- 10 Oscillographic and TV display elso use liquid crystal screens





(viii) Heat of vaporization of liquids depends upon the intermolecular today altraction present between their molecules.

(ix like does not show any vapour pressure on its surface at =1°C

(ii) Boiling point of a liquid is independent of external pressure

True (viii) True (iii) False (iv) False (iv) False (iv) True (iii) True (iii) False (iv) Flase

Q.6. (a) What type of intermolecular forces will dominate in the following liquids?
(i) Ammonta, NH₃ (ii) Octana, C₂H₁₃ (iii) Argort, Ar (iv) Proposane Anno CH₃COCH₃ (v) Methanol, CH₃OH

(I)Ammonia (NH.)

Nitrogen is a sufficiently strong electronegative element, therefore, N of one notational develop H-bond with H of another molecule of NH $_{\rm B}$. Hence in NH $_{\rm B}$ H-bondingal dominate

(lif) Octore (C_pH₁₀)

Octane is a non-polar molecule. It will not have dipole-dipole interaction or 4bonding. Its molecules will have weak London dispersion forces among themselves. (III) Argon (Ar)

Argon is noble gas and is non-polar if will have London dispersion forces arrose it molecules. Due to non-polar nature. H-bonding or dipole-dipole interactions will not bresent.

(te) Propenane (Auctone) (CH,COCH,)

It has a polar carbonyl group C=O. Due to this group, propanone will de-elliptic dipole-dipole interaction

(e) Methanol (CH₂OH)

In methanol, strong electronegative O is present. Thus O atom of one molecule will form H-bond with H of another methanol replectule

Thus, methanol chiefly, has H-bonding.

(b) Proponene (Acetane) CH₂COCH₁, proponel, CH₂CH₂CH₂OH and button. CH₂Ch CH₃ CH₃ have very startler relative molecular masses. List them in the expected of of increasing boiling point. Explain your enviour?

The boiling points will be in the following order

 Propanol
 >
 Propanone
 >
 Butane

 CH₂CH₂CH₂OH
 CH₃COCH₃
 CH₃CH₃ CH₃ CH₃
 CH₃CH₃ CH₃

 97 ° C
 56 ° C
 0 ° C

7

h is because

Limite and Spittle

- Propanol has H-bonding
- Propanone has dipole-dipole interaction
- Butane has weak London dispersion forces

Since F-bonding is stronger than dipole-dipole interaction, which in turn is stronger than London forces.

QL Explain the following with reasons.

i) Is the hydrogen banded structure of FIF, which is the strunger band: the shorter constent band or the longer hydrogen band between different stolerates?

Generally shorter bonds are stronger bond.

Further, in HF short covalent band is formed by overlap of orbitals of H and F while H-bond is just as electrostatic interaction between different molecules. Hence short covalent bond is stronger than longer H-bond.

[4] In a very cold culater the fish in gorden pends one their lives to hydrogen bonding.

In cold winter, when temperature falls celow 4°C, cold water being lighter comes to the surface. When temperature is further decreased, cold water freezes to kee Thus, an insulating layer of lice is formed above waim water. This layer of lice prevents further heat loss from tindemeath water. Thus, it saves equalic life e.g. (lithes from cold.)

(ii) Water and ethenol can mix easily and in all proportions.

Solved on Page 176

96. Erlefty consider some of the effects on our flows of if under has only very weak integers bonding present its prolocules

- In $H_{\epsilon}O$ If H-bounding is weak, it will have following effects on life.
- The freezing point of water will be lower than O°C. This will be a serious problem to the life of living things.
- Water acts as thermostat for our earth. With weater H-bonding, water will not be a thermostat. Thus, there will be large changes in the temperature of the earth.
- Due to weak H-bonding, denuity of ice will not be sess than water. Thus it would be danger for equatic life
- With weaker H-bonding, the structures of proteins and DNA would not be seablised.

ly All groups have a abstractoristic critical temperature. Allows the critical temperature it is impossible to liquidy a gas. The critical temperatures of carbon disards and



VI Serven

3

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ne are 31 14°C and 41.9°C, respectively whenter forces? Briefly explain your choice? ecty. Which gos hos the ster

The critical temperature of CO_{\parallel} is closer to room temperature (i.e. 31.1 $^{6}C)$ if shows that CO₂ is ready to be liquefied around room temperature by applying pressure. However, critical temperature of CH₄ is 81.9 °C. It shows that CH₄ must be cooled below to temperature so that it could be liquefied by applying pressure.

Since at room temperature CO_0 is easily liquefied but CH_0 not. Hence, CO_0 by stronger intermolecular forces than CH_0 at the same temperature,

Q7 Three liquids have the properties, mentioned against their names.

-	Woler	 Apparent IA cross 	Protone
Mobicular Expander	H ₂ O	C,H,O	C.H.,
Nederico antiberado mayos comos	16	58	72
Einhalpy of expectantials 🚕	41 1	34.9	277
Bothing point ("C)	100	56	36

at type of intermolecular force predominates in each liquid:

- (1) In water
- (6) In propanone
- (III) in pentang
 - H_gO has H-bonding
 - Propanone has dipote-dipote interaction.
 - Pentane has London dispersion forces.

(a) What do you deduce about the relative strength of these forces in the Hautlet half year conclusions.

The order of strength of these forces is

hil-bonding > dipote-dipote interaction.> London Dispersion forces

Hence, due to strong H-bonding b.p. of H_2O and its ΔH_2 is greater than properties which has dipole dipole interection

In pentane weak London dispersion forces are present, thanefore, its boiling point $e^{i \hat{q}}$ He are lower than both HgO and propanone.

(b) If the Squade are shoken together to pairs.

(i) Which put would be unlikely to mix?

Water & Pennage will not rube with each other since pentage is non-polar and Will in polar. Therefore they will not have attractions for each other.

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ill Exploin this immincibility in terms of the forces between the mod

H₂O and propanone are polar substances and have H-bonding and dipole-dipole interactions. Hence they attract the molecules of each other. Thus, they mix with each

However, pentane is non-polar and has weak London dispersion forces. It can not dayalop forces with water or propersone molecules, hence, it is immiscible with H₂O and епоовволе.

miChoose one of the pairs that mix and any whether the enthalpy change on mixing would be positive or negative

Water and propanone will mix with each other. This mixing is exothermic process and have we enthalpy change, because both water and acetone have strong attractions for each other during which energy is released

Q3. Describe the confuse forces responsible for keeping the particles together in the foliabing elements and compounds and their effects on physical properties positing use of the data.

Substance	Familia or	Millin Moss formal	MIP 45
Near	Ne	20	248
Argon	Ar	40	- 189
Water	H _L O	18	0
Seduce (buonice	NEF	42	993
Diamond	Ç	12	3850

- No and Ar are noble gases and have weak London dispersion forces. Since Ar is larger in sks, therefore, it has greater polarizability than Ne and therefore strong London forces. Hence its m.p. is higher than Ne.
- Water has strong H-bonding, therefore, it will have sufficiently high m.p. than Ne and Ar.
- NaF is an limit solid. It has cubic crystal structure in which each Na* Ion is surrounded by six F * ions and each F * is surrounded by six Na* ions. These ions have strong electrostade forces. This arrangement gives bardiness to NaF crystas.
- For maliting Walf crystal, large amount of energy is required. Hence, its m.p. is higher than Ne. At and H₂O.
- Diamond has network of carbon atoms, which are joined together by strong covalent bonds in a trimbedral manner
- Due to strong network of C atoms, it is very difficult to separate these atoms from each other Hence Diamond has higher m.p. than Ne, Ar, H.O and Nas



Q9. The boiling points and masses of hydrides of some first rose element are table

Substance majo	Balting Letter	Alexa Mass
CH ₄	109	16
NH ₃	240	17
H-O	373	18

Suggest reasons for the difference in their balling points in terms of the type of molecules incalsed and the nature of the forces present between them.

 $C,\,N$ and O are present in same period of periodic table

- Of these C has the least electronegative while O has the highest electronegativity
- CH_{d} is a symmetrical and non-potar motecute. Therefore, if has weak London dispersin forces intence, its boiling point is very low
- NH_b is an unsymmetrical polar molecule. In NH_b, due to high electronegativity of N,i develops H-bonding among its molecule. NHI can form one H-bond per molecule Hence, its boiling point is higher than CH4
- H₂O is an unsymmetrical polar molecule. In H₂O, strong to highest E.N. of oxygen atom. Moreover, water molecules can form two H-bonds pe

Hence, its boiling point is higher than both CH₄ and NH₅

Q10 Expidin the term naturated copour pressure. Arrange in order of increasing social pressure: $16m^3$ toster, $10cge^2$ mater, 1 dm 3 of ethanol, $50cm^3$ ethanol and $50cm^4$ ether

Vapour pressure does not depend on the amount of substance but it depends upon the pature.

Generally. Stronger the intermolecular forces lower will be the vapour pressure and vo versa. Hence order of vapour pressure will be

50 cm² of other > 1 dm² of ethanoi = 50 cm² of ethanoi > 1 dm² of $H_0O=50$ cm² of $H_0O=5$

It is because, that H₂O has strongest H-bonding and It can form two H-bond is molecule. Therefore, its vapour pressure is lowest

Chance has somewhat weak H-bonding than H₂G because ethanol can form one that does molecule. Therefore its analysis of the second of the sec band per molecule. Therefore, its vapour pressure is higher than H_eO.

Ether has only weak forces. Hence its vapour pressure is the highest-

Q11. While a volatile liquid standing in a becker suspensive, the temperature of the liquid remains the same or that of its sure of the same or that of its sure or the same o liquid remains the same as that of its surroundings. If the same liquid is allowed to upported little cames have at that of its surroundings. If the same liquid is allowed vaported little camesphere in an insulated flash, its temperature false below that of its surrounding. Explain the difference in behaviour.

When a liquid is evaporating continuously in an open container, heat from the garounding enters into the liquid Hence temperature of liquid remains equal to that of _{extroundings}.

However, if biguid is insulated from surroundings then due to evaporation its \top will fall but no heat will move from surrounding into liquid

As a result temperature of liquid falls below that of surroundings

Q12 Explain the following with reason

() Ecoporation courses cooling

Solved on Page 179

(10) Evaporation takes place et all temperatures

Faisalabad Boars, 2007: Bahawajaur Board, 2008: Gujranurda Board, 2010. Lakare Soard, 2014,

Molecules of liquids have K.E. at all temperature. Thus, they are continuously moving. They collide with one another As a result some molecules may get K.F. higher than the twenge value. Such molecules escape from the liquid surface as vapours. Hence, evaporation takes place at all temperature

At low temperature, K.E. of molecules is low, therefore, rate of evaporation is also low. However, rate of evaporation increases with increase in temperature

More or less, evaporation continuous at all temperature.

(ii) Builling needs a constant supply of bear

(D.C. Khan Board, 2009: Lahore Board, 2010)

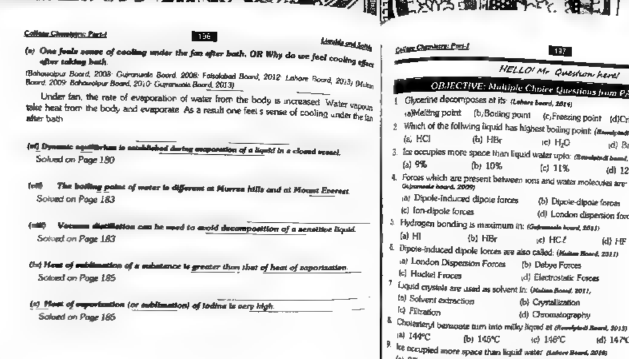
aguid holls at a temperature at which its vapour pressure becomes equal to the external DIESSUITE.

Since vapour pressure of a liquid depends on temperature, therefore, vapour pressure of a expose vapour pressure of a liquid depends on temperature, measure. Vapour pressure of a liquid becomes equal to external pressure only at a specific temperature. To seep liquid bedling, this temperature must be maintained. Thus, a constant supply of heat must be expired to the hosting figurid.

(a) Karthenware passels keep water cool

(Mukan Board, 2007 Bahawa) or Board, 2012 Labort Board, 2012 Cultimited Board, 2012) Equition again, and name amail pores, When water is placed in earthenware pots it.

When water is placed in earthenware pots it water from the whole "Notice through the ports. Due to continuous evaporation of water from the whole striage of eartherware ports, its temperature decreases. Hence, eartherware vessels lead white ports. Its temperature decreases. Whiter cool.



(a) 4.0°C

(a) 98°C

A Muree hills water boils at (Laters Seed, \$907)

(b) 100°C

OBJECTIVE: Multiple Choice Questions from PAST PAPERS a)Meliting point (b, Boiling point (c, Freezing point (d) Critical point 2 Which of the following liquid has highest boding point: (Remaphed bond, 2011) (d) Br₂ 3. See occupies more space than liquid water upto: (Remitted beaut, 201). (d) 12% Forces which are present between ions and water molecules are: us. (d) London dispersion forces (d) HF 6B 14750 9. It's occupied more space than liquid water (taken Beard, 2016) (a) 9% (d) 32% (b) 10% (c) 11% is). The boiling point of higher allower as are greater than those of lower affeates due to the Resorts that (Patestabod Board, 2010) (a) Higher alkanes have greater number of atoms. (b) The potarizeability of higher alkanes are greater (c) Higher alkansas have zigzag structure (d) Higher alkanes have greater hydrogen bonding (d) Figher alkanes have greater hydrogen transmings strongest sold emong halogen acid is (Bahasatan Baard, 1916) (a) (Jp (c) HBr (d) H (b) HCI (Vater has maximum density at (Patestabul Book, 1991)

(c) 100°C

(d) 0°C

(d) 10°C

(d, 50°C



14. The volume of the increases than liquid water by the (d) 15% let 12% (b) 9% Ç42 57%

15. Beiling point of H₂O at Mount Everest would be an o. as -4, 2010s (b) 100°C (c) 191°C æ 98°C (d) 69°C

ard, 2011)

(c) Ethane (d) Hexane

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17 The distillation of liquid under reduced pressure is called? (Sergodio Rount, 1913)

as destructive distillation (b) vacuum distillation

rd) simple distillation e transumei distillation

	Answers to Multiple Choice Questions from Past Papers.								
Q#	Aas	Q#	Ame	Q#	Ans	Q#	Ans	Q#	Ans
1	(b)	Z	εγ	-	(a)	4	(c.	5	4q.
6	[a)	7	,d)	-8	(6)	9	(a)	10	bj
11	(d)	12	(a)	13	(a)	14_	(b)	15	Id]_
16	(d)	17	(b)						

Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK!

SOBJECTIVE QUESTIONS FROM PAST PAPERS

INTERMOLECULAR AND INTRANOLECULAR FORCES, DIPOLE DIPOLE FORCES

Short Questle

Intermotecular forces are stronger than intramotecular forces. Why? 18-10

alpholi Board, \$610) (2) Define Dipole-Dipole forces and lon-dipole forces. (Res

(3) London dispersion forces are weaker than dipole - dipole forces. Why? the 2005 Off Dipole-dipole forces are stronger than Landon dispersion jorces (Mules Berl 2005)

Lorer Constitions.

1. What are intermolecular forces? Name the four types of these forces and explain any officer.

(2) What are intermolecular forces? Write the names of different types of their forces of any analysis intermolecular forces? explain instantaneous dipole-induced dipole forces. (Fetalabos Board, 2012)

(3) Define Debye forces, Amerphous solids, Allotropy and Hydrogen bonding. (Laborates)

Office Chapters Cartel IDNOOM DISPERSION FORCES, FACTORS AFFECTING LONDON DISPERSION FORCES

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(). What are London dispersion forces for interactions? mil. 207 (A

(a) How that increase of polarizability down the group in noble games is responsible for the How are increase in melting and bailing points? (0.5 Rum Bowel, 1911) OR Bailing point increases from He to Rn in VIII group. Why? (Balander Bowel, 1911) OR Bailing point increases from He to Rn in VIII group. Why? (Balander Bowel, 1911) OR Why the boiling points of noble goses increase down the group? (Dufamenter Bowel, 1914)

(ii) Melting points and boiling points of halogens increases down the group. Explain it of Board, 2009

(4) Why the molting and boding points of alkanes increase with increase in mole? hims? (Labora Board, 2007, Surgedite Board, 2009, Employed Board, 2009).

(5) Why athens (C_iFl_w is a gas whereas hexage (C_iFl_w) is a liquid at S.T.P 2 at a stam point 2010; Aunti Kashut: Board, 2019;

(1) Write a note on factors affecting London Forces. (Surpulse Seems, 2013)

PROBOGEN BONDING AND ITS APPLICATIONS

Bert Questions

20 14-bonding is present in chiaroform and acctone. Justify. Philips Barryl, 2009; D. 14-bonding is present in chiaroform and acctone. Justify. Philips Barryl, 2009.

(i) Why HT is a usering or present of the Ann HCl Statem Barrel, 2008; Lohere Bourd, 2008; D.O. When Bu 2023; Revelopmet Bourd, 2013)

(ii) Why belling point of H₀O is greater than HF? (Surguilla Bourd, 2013; Lahare Bourd, 2014)

(ii) Why belling point of H₀O is greater than HF? (Surguilla Bourd, 2013; Lahare Bourd, 2014)

Water is liquid at room temperature but H₂S is a gas. Give reason: But have been 2008.

Live Board, 2009. Geformed Board, 2011: Randplad Board, 2009, 2011. Malla Select. 2009.

Labors Board, 2009. Ordermeele Board, 2011: Raselphali Board, 2009. 1911. Praces Commun. (5) Water and ethorios can mix in all proportions, give rection. (Labors Board, 2018; Cal Lower Scale) of the rest ranks are insoluble in water Why? (Labors 10).

(7) Write dawn two applications of hydrogen bonding (8)

(B) Power communo applications of nyarogen persungs institly the statement, (substitute compounds are soluble in a polar solvents. Justify the statement, (substitute compounds)

the floats on water Justify it OR fee occupies more space then water give reason. OR How see life survives under frozen and OR Density of low is uses than liquid water brillion water than the state of the survives under frozen and OR Density of low is used than liquid water brillion water than the state of the Explain with region, Génica Board, 2087; Februinal Board, 1908, 2011; 2013; Labora Board, 2009; Sagodho Board, 2008; Ballion Board, 2009; Especial Board, 2010; Enceptival Board, 2010; D.G. Kiew Board, 2010; Enceptival Board, 2010; Enceptival Board, 2010; Enceptival Board, 2010; D.G. Kiew Board, 2010; Enceptival Board, 20

Water freezes from sturface to the down direction in ponds and lakes. Explain with Peason, (Behavious Bount, 2017) OR Lower density of los than water how got significance

Comment (Newton Board, 2008) aı, Water freezes from surface to the dounward direction in pands and falses. Explain tahy? (D.O. Hhom Bound, 2014)

fem directors:

Define hydrogen bonding. How dose it exploin the indicated properties of following statements. Exhaus Sound, 2010)



Annale and Spins

- (2) Explain hydrogen broading in NHs, HsO and HF, how it is helpful in explaining the structure of 10s. (Labora Board, \$612)
- (3) What is hydrogen bonding, Explain role of hydrogen bonding in food and biological material diagodia Board, 2011.
- (4) How does hydrogen bonding explain the formation of ice and its lesser density than figuid water (Gujramente Board, 2012)

EVAPORATION VAPOUR PRESSURE

Short Question

- (1) How the rate of exaporation depends on the surface area? (Lakore Board, 2009)
- (2) Looporation of a liquid course cooling, Expirin. D.C. Khan Beard, 2007. 2009: Nature & 3010: D.G. Khan Seard, 201F. Surpetite Board, 2007. 2014: Friendschad Board, 2009: Labor: B. 2019. 2011. 2014.
- (3) Define evaporation and vapour pressure. (Felestated Board, 2017) Ans: Take defined
- (4) Define vapour pressure of a liquid. (Fateriahed Beard, 2009)
- Define vapour pressure Write down two factors that affect vapour pressure of a liquid. (Remelpind Board, 2012)
- (6) Why the vapour pressure of water Ethyl alaphoi and diethyl ether are different from each other at O'C (European Sound, 2007)
- (1) Define Vapour pressure. Write a method for determination of vapour pressure of a
- figuid: (Bahmadper Boerd, 2009 Statem Board, 2015: Eurgadia: Board, 2013, 2014)

 (2) How suppour pressure is measured by Manameter method. (Fairolabed Board, 2007: Gajrannaia Board, 2009; 2011: D.G. Khain Board, 2012; Midden Board, 2012)

BOILING POINT, EFFECT OF EXTERNAL PRESSURE ON BOILING POINT

Short Questions

- (1, Define bolling point translated Board, 2015) OR Define boiling point. How it is affected by 60767903 Overstare? (Remoteball Roans, 2010)
- (2) The bolling point of water is different at Murree Hills and at Mount Everst. Explain with reason. (Felicational Bourd, 2007, 2010; Bergodine Bourd, 2013) OR Water bolls at loss bemperature on Hills, areas than on plans preces. Why? (Behavelore Bourd, 2010)
- (3) Explain the working of pressure cooker (Formation Board, 1912, OR Food can be cooked quickly in Pressure Cooker Exploin with reason, (Surpollia Board, 2011-2014)
- (4) Vacuum distillation can be used to avoid decomposition of a sensitive liquid Explain Bahasahar Board, 2012: Labora Board, 2012; OS What Is the Importance of working later (Superments Board, 2009, 2013)
- (5) Why different liquids possess different boiling points? (Mailen Board, 2013)

- 1) What is boiling point? What is the effect of external pressure on boiling point? Why the temperature of a limit of the state of the temperature, of a liquid remains constant at boiling point although heat is continuous; supplied? Mohore Board, 2018)
- (2) What is the effect of external pressure on boiling point of a substance? Give extend that were 2012;

20+ HOLAR HEATS OF FUSION, VAPORIZATION, SUBLIMATION

Short Questions

- (1) What are motor heat of fusion and motor heat of sublimation. (Sergodia Based, 2016)
- (1) What the motor heat of Japanisatian and motor heat of fusion (General Board, 2010)
- (3) Define molar heat of vaporazation with one example. (Remotive Board, 2009)
- Define from rease by supersecuting user one example. Reasely, 2007.
 Why heat of sublimation of a substance is greater than that of heat of vaporization forgodie Book, 2013; Federal Book, 2014; Federal Book, 2015; Federal Book, 2016; Federal
- Readphold Board, 2010: Lancer Board, 2012: Department Board. 2012: 2015; Lahors Rourd R. (6) Gasoline evaporates much jaster than water Give reason. (Outcommise Board, 2014)
- (7) Why the heat of vaporization of water is greater than that of CH₄ (0.6 10cm Board, 2011)
- (5) Why heat of vaporization of hearne is greater than that of ethane (8-repella Bound, 2007)
- (9) Steam couses mans severe burns than does the boiling water (Garanada Band, 2010)

LIQUID CRYSTALS AND THEIR APPLICATIONS

- Bort Questions
 (1) How liquid crystals act as temperature sensors? (Labore Board, 2007-Holom Board, 2013)
- (2) How liquid crystals are used as a locator of turnors? (D.G. Rhom Board, 2009)
- (4) The High Carponine are used as a recursor of surrors. 1979. Been \$1907. Housiphak Bound, 2809. Multin Bound, 2012. Labora Bound, 2013. 1979. D.O. When Bound, 2014. Falsokabed Bound, 2013. Surpodha Bound, 2014. Amer. All uses are given below. Generally, two or four uses are asked in

Onemone

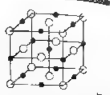
- (I) What are liquid crystals? Give their two uses (or three uses) in daily life. (Falantabad Sound, 1909, 387). Departured Sound, 2016; D.G. Khan Bound, 2009; Ased Rashmir Bound, 2012; Latere Board, 2010; Control of the control of
- (2) Glise four uses of liquid crystals, (Unbarrenta Board, 2008, Emposite the
- Series pour trasses y requisit triposters component uses of liquid crystals (Germedia Series, 2012). Gipermental States, 2013: Labora Beard, 2014.



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Library and and



PSTRODUCTION

Exercise Q.4

What are solids? Give general properties of solids. How do you differentiate between crystalkine solids and amorphous solids?

Solid

It is the etate of matter, which has the strongest intermolecular forces, definite shaps and definite column

In solids atoms, ions and molecules are held together by strong ochesive forces and these cannot move at random

Most of the solids exist in crystal form

Figure (1) Studies (Classification of solids on the basis of <u>Recolutive of Portfoles</u>)

On the basis of regularity of solid particles, these are divided into two types 1. Crystalline souds 2. Amorphous solids

Oysedfine stands

The solids in which atoms, lone or molecules hose regular, three-dimensional errorgament are called crystalline solids.

NeCl, Diamond etc

2. Antorphony of or

The solids in which stome lone, or molecules do not have a regular ariest errangement are called amorphous solid.

These are called oscudo solids.

Properties.

- 1 These bave rendem structure.
- 2 These have short range order like liquids.
- 3. They don't posses repetitive structure of crystals.
- 4 They may have hardness and desticity

College Chambers, Pers.

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South and Cate

- 5 They do not posses sharp melting point
- 6. These have indefinite heats of fusion.

Expresion:

Glass, Rubber Plastics, Glue etc.

NOTE:

- Operatifies In amorphous solids long large order is absent. These solids have areas of short-range order in their structure. These areas are called crystallites.
- Super-Cooled Lightder Many cryateline solids can be changed into amorphous solids by melting and then cooling rapidly. In this way, particles can not arrange themselves regularly. Thus, they have definite volume and shape but arrangement of particles is not regular. Hence these are also called super cooled liquids.

Difference between Crystollins and Amorphous Solids

Crestalline Solide

These have definite geometric shape in which atoms, ions or molecules have a regular and repetitive three dimensional arrangement

These have sharp melting points.

These have definite heats of histon

Examples, NaC., diamond, KCi etc.

These are anisotropic

These are called true solids

solids 5

Amountmen Solints.
These do not have definite geometric shape in these atoms cons,or molecules have a condor or non-repetitive three dimensional arrangement.

These do not have sharp melting points.

These have indefinite heats of fusion.

These are isotropic and resemble liquids

These are called pseudo solids.

Examples Glass Rubber etc.

Ersycles Q.6 (a):

Define unit cell What are unit cell dimensions? How the idea of crystal tattice is developed from the concept of unit cell?

The smallest part of the crystal lattice, showing all the characteristic features of the whole crystal is called a unit cell.

But Coll January and a Cognollary and a Departs

A unit cell is identified by the sength of its faces i.e., sides 'unit cell tengths) and the angles between these faces (unit cell angles).

There are three unit cell lengths represented by $a,\,b,\,$ c and three unit cell angles denoted by $a,\,\beta,\,\gamma$ as shown in floure.

UNA COU



Angle between sides "b" and "c" is "a"

Angle between sides 'a' and 'c' is '\$'.

Angle between sides 'e' and 'b' is 'y'

All the six parameters of a unit cell are called unit cell dimensions or crystallographic

- Thus complete information of crystal is present in its unit cell.
- If structure of a unit cell is snown, then structure of whole crystal can be made by repeating it.

These are seven different types of unit cells depending on the rength of faces and angles So there are seven types of crystal lattice/crystal systems.

It to an array of points, representing atoms, tons or molecules of a creat different oldes in three-dimensional space.

In crystals, particles have definite positions. These are separamented by points in a crystal. These points are called lattice goints or whice size. This three dimensional arrangament of points in a crystal is called crystal inition or mace lattice, e.g., NaCl



Triport Lives Tripoglation

Exercise Q.6 (b):

Lapton seven crystal systems and draw the shapes of their unit cells

CRYSTALS AND THERE CASSIFICATION

The group of crystals whose shoper are built up by only one type of unit cells a celled crystal systems.

There are seven different types of courtal systems.

Lobb System

All the sides are equal

All the angles are of 90°

a=β=γ=90*

Examples

Fe. Cu. Ag. Au, NeCl. NaBr diamond etc.



Color Charleton Part I

To transmit Sestem Out of these, two sides are equal All the angles are of 90°

a=0=y=90"

Employee

So, SnO₁ MnO₂ NH₄Br etc.

\$ Gritan Countrie Severes

All the sides are unequal axbec.

All the angles are of 90° **π=β=γ=90°**

to make

indine, Rhombic aulphus, K₂SO₄ BaSO₄ etc

6. Inwend Or Broadfule and Sestera

All the order are equal

a = d = 6

All the angles for between 90° and 120°

1=0=y=90°

E TOTAL

Bl. AuO, NaNO, 1010, etc

thesiopsical System.

Out of three, two sides are equal

 $^{
m T_{MO}}$ angles are of 90° and third is of 120°

 $\alpha = \beta = 90^{\circ}$ and $\gamma = 120^{\circ}$

Graphile, ZnO. CdS, loe Zn, Cd etc.















20%

College Chembers: Ports

Montu linte Nestem

All the sides are unequal ambec Two angles are of 90° and third one is greater than 90° $\alpha = \gamma = 90^{\circ}$ and $\beta > 90^{\circ}$



Sugar Monoclinic Sulphur, Na₆SO₆, 10 H₆O, Borax etc

7 Inclinir System

All the sides are unequal a + b + c All the angles are unequal a # B # Y = 90°

Exemples

H,BO, CuSO, 5H,O, K,Cr,O, etc



Licente and Sally

		1,0003	(78) a cosof \$	stants
St. 1	Constant Byrolotta	Aus	Angles	Examples
1	Cubic	a=b=c	¢=β∞γ≠90°	Fig. Ca., Ag., NaCl., NaSk. Diserround
2.	Tetragonal	amb≠c	a=\$=r=90°	Sn. SnD _p , MnO ₂ , NH ₄ B ₁
3.	Orthorhornios	a≠b≠c	gc=β= γ≔90°	lodine, Rhombic sulphur 8650,
4.	Rhombohedral or Trigonal	ambec	a + 8 = y + 90°	EL, ALO, NANO, KNO
5.	Hesagonal	n≖b≠c	n=8=90°,+- 120°	Graphite, ZeO, CdS, Jor, Zn, Cd
6.	Manoduric	n*b*c	0=y= 40°\$ ≠93°	Sugar, Sulphur Bores, Ne SO ₁ 10H ₂ O
72	Tricknet	*b#c	a###7#90°	H ₁ BO _p , K ₂ Ct ₁ O _p , CuSO ₄ SH ₃ O

Explain the following properties of crystalline solids. Give three examples in early case (iu) isomorphism

(Bi) Habit of a crystal (uli)Polymorphism (i) Anisotropy (ii) Cleavage (v) Transition temperature (vi) Symmetry (vi) Growing of a crystal

PROPERTIES OF CRYSTALLINE SULIDS

Colors Chambers: Part-J

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and Salvile ووايوني

beat wifeed shape Costalline solids have definite, regular three dimensional arrangement of particles.

Each crystal has definite faces (sides) and definite angles interfacial angles. between the laces

For a given crystal, the interfactal angles, at which the surfaces intersect are always the same no matter in which shape they are grown. The locas and angles of crystals are not changed even if it is ground to a fine powder

2. Meldin porti

Crustalline solids have definite melting point and can be identified by this. Thus crystels have definite heats of fusion.

3. Clemente and Clemonge Plane

The breaking of crystalkine solids into smaller (dentical crystals due to external presents is called cleavage.

The plane in which a crystalline solid can be broken into smaller identical crystals is ulid cleavage plane.

The cleavage planes are inclined to one another at a particular angle for a crystalline solid. The angle is different for different solids

Actsorphys

Some of the crystals shall parieties in physical properties depending upon the direction. Such properties are called adoptropic properties and the phenomenon is called autocolled anisotropy.

A substance, which has this property is called an anisotropic

Crystals show the property of anisotropy. It is because crystals have different arrangement of particles in different directions.

Eggpt es:

In crystals, electrical conductivity thermal conductivity, co-efficient of thermal

expension and refractive index are anisotropic properties

Electrical conductivity of graphite is greater in one direction than other it is

Electrical conductivity of graphite is greater in one direction than other it is

because in graphite mobile electrons can move easily parallel to its layers rather than perpendicular to its layer

Cleanage itself is an anisotropic behaviour

The properties, which do not depend upon the direction, are called isotropic Properties, e.g. melting point dic

The substance which posses isotropic property is called isotrope.

Scharnerre

estimates.

The reportition of faces, angles and edges when a created to rotated by 360° along its City is collect symmetry.



Crystels have vertous types of symmetry elements e.g. centre of symmetry, plant symmetry was of symmetry sto

The alongs in which a wyotal woully group to called habit of a crystal.

e.g., where a saturated anisition of NaCl is cooled, cubic crystals of NaCl are formed Thus NaCl hea cubic hebit

Caystais are obtained either by cooling its seturated solution or by slow cooling a liquid substances. These are formed by growing in various directions in patible conditions, shape of growing crystal is not changed. However, if conditions are shape then shape of the crystal may also change

e.g. Cubic crystal of NaCl becomes needle like when 10% ures is present in soluti as impurity

2. Ison organism

The physicalities to collect incomplism.

These different substances are called isomorphs of each other.

- Isomorphous substances have different physical and chemical properties.
- The formula of isomosphous substances shows that they generally have same used
- A crustal structure depends only on the number of atoms and then wall combination, it does not depend on chemical nature of atoms
- Somorphous substances crystalize together in all proportions in a homestics mixture

Examples.

	4000	St. Co. H
NeNO, KNO	Enomobol admi	1 1 3
K,SO, K,CrO,	Oytharhombic	2 1 4
ZrSO _e NISO _e	Odhodombie	1 1 4
NaF MgQ	Cubic	1 L

- The structure of negative ions like NO_6 and CO_8^{4n} are same Both are Marsh stance.
- glanux. Similarly the Sinucture of $SO_4^{\,2a}$ and $CrO_4^{\,2a}$ are also some. Both are introduced

1. It Is morphism

- 20% The substance, which exists in more than one crustalling form, 19 gailed polymorphic,
- Officers crystalling forms of a substance are called polymorphs.
- Polymorphs of a substance have some chemical properties
- Polymorphs have different physical properties, it is due to different structural arrangement of their pertices.

- AgNO, exists as Rhombohedral. Orthorhombic
- CaCO₂ exacts as Trigonal and orthorhombic

Difference between Isomorphism and Polymarphism

Isanimphism	Professionalities
Different substances exist in same crystalline forms.	Same aubstance exists in two crystalling form Polymorphic forms have same
different physical and chemical	chemical properties but different physical properties Polymorphic forms can be inter-
Isomorphism substances cannot be inter-converted Isomorphic forms have different	converted forms have same formula unit
Formule units Examples Both MgO and NaF are 5	Exempted CacOh endsts in trigonal and orthochombic form
	Different substances exist in same crystalline forms. some crystalline forms. isomorpi rous substances have different physical and chemical a properties. Isomorphius substances cannot be inter-converted. Isomorphic forms have different formula units.

tition in s

rs in which as element axists in more than one crysta

Different custalline forms of an element are called allowards or ellotropic forms Tropies:

- Carbon can exist in graphite (Flaxegonal) and diamond , cubic) forms
- Sulphur can exist in Rhombic and Manachinic forms
- Tin (Sn) can exist as gray an(cubic) and walls sin(tetragonal)

lighter retires from personale The temperature at which per organization forms of the so discuss with such other is colled transition temperature



Example Sulpher whombir I 11C-Grey Time enduce

KNE) wather hombic NegSD₄ 10ht₈O(nymeted form)

32 38 G Na_BCO_B.10H_BOthigher hydrated form) 3

Links and Sale

Sulpher , monoclinic i White Tirt (Telingonal

KNO_s(rhomboheshall)

NegSO_{st}anhydrous term) + 10H₂O Na₂CO₃ 7H₂Otlower hydroted form. +3H₂O

At this temperature one crystalline form of a substance changes into other

int.

- Above and below this temperature, only one form cuists
- The transition temperature of the allotropic form of an element is always sea thans melting point

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THEATON OF LONG

(On the basis of forces present in the crystal)

Crystal have been classified into four types on the basis of forces present between app ions or molecules in them. These torces are also called coherive forces. These torces way or chemical bonds or some other interactions.

There are four types of crystalline solids on the basis of nature of forces present to then

- A. loruc solids
- B Covalent solids
- C. Molecular solids
- D. Metallic solids

Expreher Q.7 (a)

What are same solids? Give their properties, Explain the structure of NaCi. Sketch to the to justify that unit cell of NoCt has four formula units in it.

The crystolline solids, in which positively and negatively charged tone are held together the descendants forces of attractions, are called lonic solids. These forces are also often

Examples.

NaCl. BaCl₂ K₂SO₄ etc.

Properties of Land Sofuls

- These are <u>crystalline solids at soom temperature</u> due to definite arrangument of out
- Under normal temperature and pressure, these never exist as liquids or gases.
- 3. These are <u>stable compounds</u> in ionic crystals, ions are held together by significant contests forces, which are difficulty to the contests of the contests electrostatic forces, which are difficult to break. Hence these are stable

- 4. These have high meiting point and boiling point due to strong electrostatic forces
- 5. These have key volatility due to strong electrostrile forces
- 6. These are very hard. It is because, in these crystals fonic fonces are present which are non-directional. As a result ions are attracted from all sides, Hence ions are closely packed and there is not much free space in these crustals
- 7 The structure of an ionic crystal depends upon radius ratio. <u>[onic crystals having same</u> radius ratio have same seconeiro.
 - eg, both NaCl and CsF have same radius ratio hence they have same geometry i.e.
- 8 lank crystals do not exist as individual, neutral, independent molecules. Therefore term formula mass is used to express their mass instead of Molecular mass.
- ionic crystals do not conduct ejectricity in solid state. Because in solid form, ions have fixed position and therefore can not conduct electricity

However these conduct ejectricity in molten state or in agreeous solution. Because in these torms, crystal lattice are broken up and ions become free. These free ions conduct

20. ionic crystals are highly bottle.

If it's because, tonic crystal consists of parallel layers in which cations and anions are present in alternate positions. Thus, when a force is applied on the crystal one





layer of uses slides a little bit over the other layer. In this way like usus come in front of each other

Which repet each other and thus a crystal is broken and show brittleness.

- U. Due to close packing of tone in a crystal, these have high densities.
- 2 These are soluble in polar solvents like H_gO and produce with
- 13. Since these crystals produce ions on dissolving in polar solvents. Thus their <u>reactions in</u> polar solvents are very test.
- 14. They show isomorphism and polymorphism.

Continue of Notal Cristal

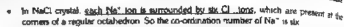
- NaCi crystal is face-centred cubic and consists of Na* and CI* ions.
- Na* has 10 electrons, while CI has 18 electrons Trus CI is After in size than Na ' ion.







Literate and Sales



 Similarly each C1 upn is also surrounded by six Na" sons. Its co-ordination number is also six.

Calculation of number of No' and Cl' lone per unit cell. Number of Cl' toru

The figure shows that in NaCl cyretal two types of Cl* ions are present

- (i) 8 CL tons are persent at the corners Each of these is shared by 8 unit cells. Hence share of 1 unit cell = $\frac{8}{8} = 1$
- (ii) 6 CI form are present at the faces. Each of these is shared by 2 unit cells. Hence share of 1 unit cell = $\frac{6}{2}$ = 3.

Hence one unit off has 3 + 1 = 4 Cl sone

Mumber of Ne* tons

The figure shows that in NaCl cyretal two types of Na* ions are present.

- If 12 Na* tons are present along the edges, each of which is shared by four unit of. Hence share of one unit cell = $\frac{12}{4}$ = 3
- (ii) 1 Na* son in present at the centre, which is shared only by one unit cell. Hence one unit cell has 3 + 1 = 4 Na* ton.

Total number of No* and Ci* lone

One unit cell of NaCl has 4 Na* ions and 4 Cl. jons. Thus the ratio of Na* and Cl. jone is 1 . L.

Facts about NaCI:

- In NeCl, the distance between two similar ions is 5.63 A*.
- . Thus the distance between two different ions is $\frac{5.63}{2} = 2.815 \text{ A}^{\text{b}}$
- In NaCl crystal, no independent NaCl molecule is present. However, NaCl rackfills
 exist in vapour phase.
- The radius ratio of NaCl is <u>0.53</u>

Color Chestern Perts

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Limite and South

Smrs Energy

fire amount of energy released when one male of the laste crustel is formed from the passeous tons in called lattice energy.

In this case the energy is given negative value

The energy required to break one male of solid into isolated loss in the gas phase is called lattice energy.

In this case the energy is given positive value. Lattice energy in instantived in $k_0 \; {\rm mpc})^{-1}$

Examples.

Nature + Cities + NeCles

ΔH=-787 № mole⁻¹

 $NaCl_{ex}$ \rightarrow $Na^*_{ex} + CC_{ex}$ $\Delta H = +787 \text{ kJ mole}^{-1}$

Lattice energy decreases with increase in size of entions or anions

It is because due to increase in size, the padding of sons becauses less tight. Thus, more empty spaces are present and structure is loose. So, less amount of energy is required to break it. Hence, radice energy of crystals decreases with increase in size of cations or anions.

Exercise Q.7 (b):

What are consider solids? Give their properties. Explain the structure of diamond

OVALENT SOLIDS

The crystalline solids in which neutral atoms are held together by covalent bands are alled covalent argencies

- These are also known as atomic solids.
- The atoms may be of same or different elements.

Covalent crystals are of two types

- (i) Glant Copalant Solida: The crystal in which atoms are joined together to form by molecules.
 - Examples, Diamond, SiC or AIN etc.
- (4) Layered Covalent Solids: The crystal in which atoms are joined together to form separate jayers.
- Examples graphits, codmium rodids and BN etc



ntaka sa Comaligna Salabag

Tree . stars some a transformational reduced of stores somet logate ; considerat percents.

- 2 Space conservation are forecast band, therefore, considerable feet space. present in these waters want their packing is located from norm or said County
- There are ben't and a so of energy a required to break them.
- These time tight melling source and bodies points.
- These stave one intention, due to strong covalient bonds
- to be these to feet out are preparal, hence these are generally bed continues to CONTROL

Homewer an graphite pertalled layers have free electrons. Hence it is a good conducts of electronity penalties to its ages. However, graphite is not a conductor perpendicular to its invers

- These are <u>rachable to polar solvents</u> the water but WANTER TOTOLOGIC WHISTIS THE BETTERNE COLL ex-Covalent orystals having big molecules like dismond. SrC etc are insoluble in all solvents, it is because they cannot develop forces with solvers molecules due to their big eize
- 8 These crystal show very slow reactions.

Secretary of Degreend

Diamond is an allotropic form of carbon

There are four valence electrons in carbon.

The four orbitals (one 2s and three 2p) undergo sp⁸ hybridization to give four sp³ hybrid orbitats.

Thus, each carbon has four and hubrid orbitals, which are directed towards by comers of a regular tetrahedron. This is the unit cell of diamond. Many unit cells undergo go overlapping to form a huge threst dimensional structure of diamond go go overlapping form covalent. bonda

Each carbon is unked to four other carbon atoms through covalent bonds. The structure of diamond is continuous and carboncarbon overlapping form a big structure. The whole crystal looks like a huge carbon molecule. Thus, diamond is also

In diamond all the bond angles are of 109.5° and the bond lengths are 154.0°. The overall structure of diamond of overall structure of diamond is face control oubic lattice.

Carlotte Charles Con !

Emicter Q.7 (c) What are malecular crystals? Give their properties

MOFECULAR SOLIDS

The crystalline solids in which point or non-polar atform the crystal are culted molecular crystals

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e.g. In solidified noble gases, non-polar atoms are present

Two types of intermolecular forces are present at molecular solids

- a. Depose-dipole inseraction.
- b. van der Waal's forces

These torces are man in weaker than some or covalent bonds

Exemples

lce Sugar Pour Molecular Crystals:

Non-Polar Moleculer Crystals, sodine sulphur P CO2 etc

Properties of undeenlar Solids

- These have request attangenent of atoms in molecules. The position of atoms can be determined by X-rays analysis
- These are soft and easily compressible due to the presence of week intermolecular forces.
- These have low melting points and boiling points due to the presence of weak intermolecular forces
- These are mostly <u>votatile</u> due to the presence of weak intermolecular forces.
- These are bad conductor of electricity, due to the absence of ions or free electrons.
- These have large errors spaces, in a crystal hence these have jour densities.
- Sometimes, these are transparent to light.
- B. Polag molecular crystals are spluble in polar solversts, while pon-polar molecular crystals are soluble in non-polar solvents.
- 9 Generally, polar molecular solids have higher melting points and boiling points then non-polar molecular solida

Stewart of L

In solid form, I₃ molecules have a layered structure In solid is, each ! . I bond distances 271.5 pm This distance is

much langer than I distance in gassous todine, which is 250.6 pm. This structure shows that I is a poor conductor of electricity. of electricity





Exercise O.S.tel.

Give different theories of a metallic bond. How does electron sea theory sustifies the electron conductivity, thermal conductivity and shining surface of metals?

METALLIC SOLIDS

The metal atoms to a metallic crystal are hold together by special type of bonds called allie bonds

Theraies of Mounting Bourling

To explain metallic bonding, following theories have been proposed

- 1 Electron gas theory 2 Valence bond theory
- 3 Molecular orbital theory

I lecture Gos Theory.

It was proposed by Drude and Improved by Loren (1923),

According to this theory, all the atoms of the metallic crystal loose their valence electrons These electrons form an electron pool or electron gas in which positively charged nuclei at present at definite positions at a measurable distance

The electrons of the sea are not attached to a particular nucleus. These electrons are delocalized over the entire crostals. therefore, these are called Free electrons,

The positively charged nuclei are held together by the free electrons throughout the lattice. The force, which binds a metal cation to a number of electrons around it, is called Metallic



2 Unleave Bread Theory

It was proposed by L. Pauling

According to this theory, metallic bonds are actually covalent bonds. However, these covalent bonds are biobly delocated and extended over whole crystal.

Molecular Orbital Theory for Band Theory)

According to this theory, valence orbitals of metals overlap with each other to feet delocation orbitals. These are called molecular orbitals. Molecular orbitals are spend over the whole crystal. The electrons in filled orbitals are localized

Actually, when argo number of valence orbitals of atoms overlap, they produce logic number of molecular orbitals. These molecular orbitals have very close energy stales and thus form a based of communication of the comm thus form a bend of energy states. Hence, this theory is also called Band theory

The energy difference between two bands determines the properties of metals crustals

College Chemistre, Part-I

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Limits and Solids

powerlies of Mensitic Solids

owthom of propopulation on the busin of Electron Gas Theoret

L. Hertifeol Conductivity

Metals are good conductor of electricity. When a metal is connected between two poles of battery, the mobile electrons begin to move towards positive pole and the electrons enter into the metal from negative pole. In this way metals conduct electricity as shown in the



Sometimes electrical conductivity decreases with increase to territière fure

It is because, increase in temperature, increases the vibrational motion of positive nuclei. These nuclei produce hindrance in the motion of free electrons.

Hence electrical conductivity decreases

2. Thermal Conductions

These are good conductor of heat.

During this free electrons take up heat from one end and transler if throughout the crystal during their motion and collision with other electrons. Thus free electrons cause thousal conduction in metals

Lestons Surjace

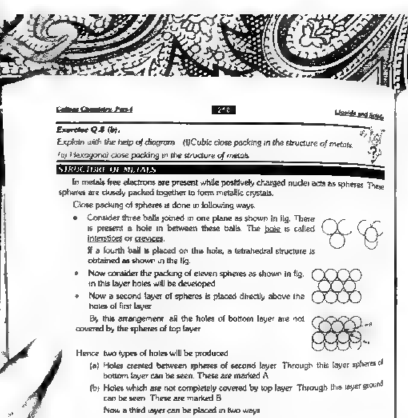
Melals have justious surface

The freshly cut surface of metals show shiring (fustrous surface). It is because when a light strike the tree electrons in the surface they are excited. When these excited electrons come to their original position, they emil light. Thus light appears to be reflected by metal surface and It appears shiring.

Multiplide And theirthe

Metals are maileable and ductile. When a force is applied on metals, their layer ship over each other and their shape is changed. Hance, these can be converted into sheets (mateable) or wires (ductile) without breaking.





it third layer is placed directly above the holes marked A, then the spheres of

If third layer is placed directly above the holes marked B, then the spherel of all the figure transmit will have different and because the spherel of the s

therd layer will be directly above the spheres of first layer

This structure is called hexagonal close-backed structure.

 This arrangement produces an ABCABC or 123123 patiern. This structure is called lace centred cubic lattice.

This arrangement produces an ABAB or 1212 pattern

the three layers will have different pottern.

(if the organist flow Probed Structure)

- (1 -)



719





Type of Solid	Simple Particles	Fattie: Types of Interterologistic Frances	Solats Typical Properties	Examples
Metallic	Cations plus delocations electrons	Metalic bonds	despites to the front of its very hard, my land pour if spire from low to vary high. furtions could, makeable, very good conduction of hear and electricity.	Na. Mg. Ai Fiti Zhi Ciri Ag W
tonic.	ations and ntilibra	Electronicia. attraction	form, moderate to very high melting points, non- condy rism of whichtaby four good electrical conduction in the motion state:	MaCI: rounds MgC
Melroules	Molecules (stars)	scendon and or dipole-ripole and, or hydrogen knock	cole, low analting pointly grow-conductions of Sense and electricity solution enable in month coars	Notice gas stements. CH ₄ (410 ₁ , P ₄ , S ₃ , F ₄ , F ₅ O
Metwork c evaluati	Aunera	Covalent bonds	yesy hard, very high pariting points non-conduction of electricity	C(damena)) SIC SIC ₁

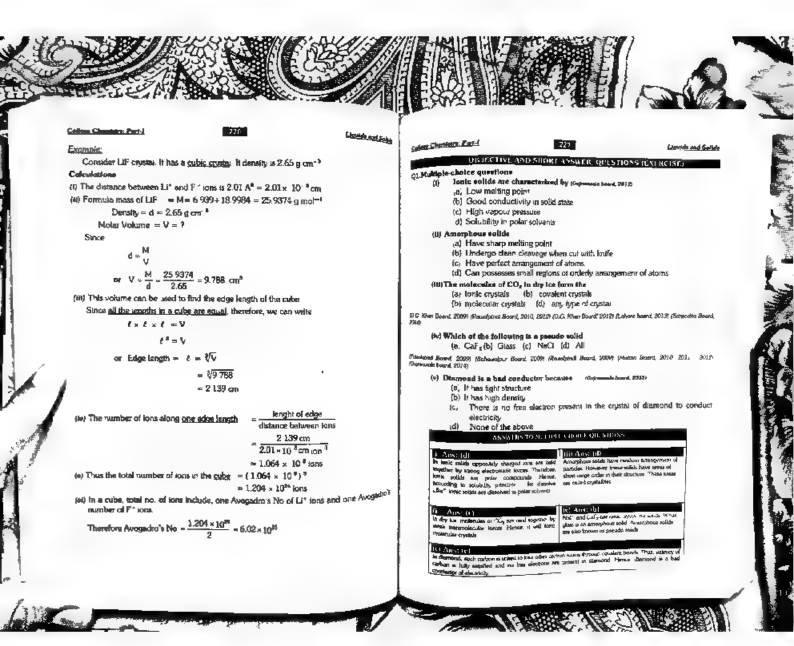
Strong arms for a transfer or attition

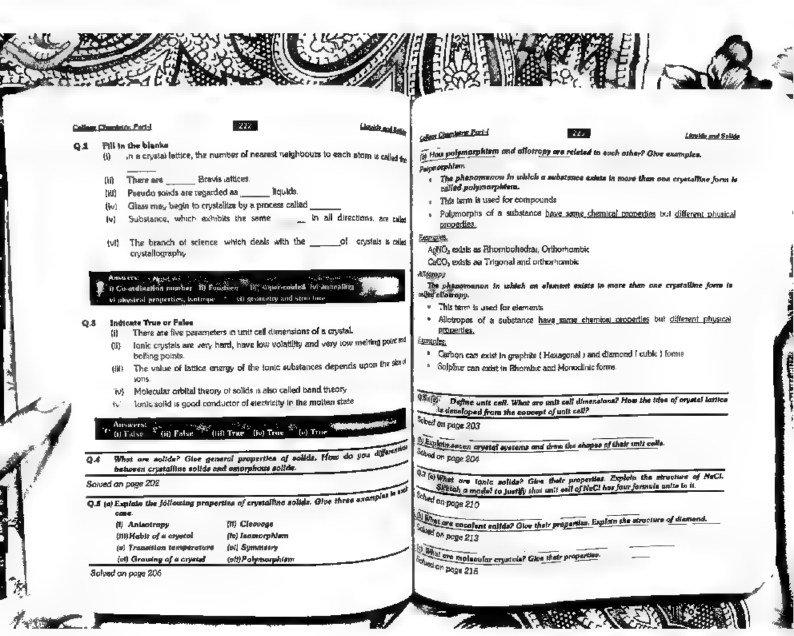
Marty methods are used for the calculation of Avogadro's Number

An accurate method is based on the study of crystab

To determine N_{α} by expitallographic method following data is required

- The volume of one mole of a crystalline solid.
 - It can be determined from its density
- The distance between particles in the crystal lattice. If can be determined by X-rays







College Chamistre: Part-I

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Linds entitle

(d) Justily that molecular erpstals are softer than lonic crystals.

(D.G. Khan Board 2009)

lonic solids are very hard. It is because these crystals have non-directional lonic tord Jonic solids are year many in a december Hence ions are closely packed and there say much free space in these crystals. Therefore, these are hard.

But molecular solids have very weak intermolecular forces. Thus these are soft an easily compressible

Q.S (a) Give different theories of a metallic bond. How does electron sea theory jurish the electrical conductivity, thermal conductivity and shining surface of metals?

Solved or page 216

- (b) Explain with the help of diagram
 - (i) Cubic close packing in the structure of metals.
 - (II) Hexagonal close pucking in the structure of metals.

Solved on page 218

Crystals of solts frocture easily, but metals are deformed under stress with frocturing. Explain the difference.

Saits are tonic and tonic crystals are highly brittle and can fracture easily.

It is because, fonit crystals consist of parallel layers in which cations and anions are present in atternate positions. Thus, when a force is applied on the crystal, one layer of ions slides a little bit over the other



layer. In this way like ions come in front of each other which repel each other and this

crustal is broken. However metals do not consist of oppositely charged ions. Thus, when a force is apply the second of the second oppositely charged ions. on metals, their wyers just slip over each other. There is no repulsion between the life. Hence these can be converted into sheets (malleable) or wires (ductile) without breaking.

Q.10 What is the coordination number of an ion? What is the coordination the cotion in (a) the NaCl structure and (b) the CoCl structure? Explain

remon for this difference? The number of necress neighbours of an ion in an ionic crystal is called co-ordinate.

e.g. In NaCl crystal, each Na* ion is aumounded by six Cl* tons, which are present sufficient for the company of the company o corners of a regular octahedron. So the co-ordination no. of Na* is six.

Similarly each Ci " ion is also surrounded by six Na" ions. Its co-ordination no. is also

- In NaCl, the co-ordination no. of cation i.e. Na* $_{\rm HS}\,\delta$ el.
- In CsCI the co-ordination no of cation i.e. Cs* is 8 ы For this difference see Q.12 , part xv

QIII Give examples of lasic solids, molecular solids and covolent macromolecular salids. What are the factors which determine whether each of these types of solid will dissolve in water or not?

The gangral solubility principle is "Like Diamotre Like" Thus, polar substances dissolve solvents and non-polar substances dissolves in non-polar solvents. Water is a polar schools Following factors affect the solubility of substances in water.

- (this large Polar substances dissolve in water easily.
- (ii) Leating Energy: Higher is the lattice energy, difficult is the crystal lattice to break, persons, lesser is the solubility.
- (iii) Sise of fons/molecules: The solubility of a substance in water decreases with increase in size of ions/molecules

logië Solide

Examples: NaCl, BaCl, , KeSO, etc

These are polar therefore; generally soluble in water. However, kinds solids with high lating emergy, or range size ions are less soluble, e.g., NaF (High lattice energy) and BaCO. (Lange size soms)

Nélecular Solida:

Mining fea:

Polyrennierular solids: Ice, Sugar

Nan-Polar molecular solids: todine, sulphur, P., CO, etc.

Polat molecular solids are generally soluble in water, while non-polar molecular solids

e solide in non-polar solvenis Cotalent Matromolecular Solids:

Empires: Diamond, StC etc.

Covalent solids are non-polar so these are generally insoluble in polar solvents like water but solublish non-polar solvents like bansens, CCL, etc. However, covalent macromolecular tolds like diamond. SiC etc are insoluble in all solvents. It is because they cannot develop force uses. lottes will solvent molecules due to their big size

0.12 Explain the following saith receous:

226 Lieute and Son College Chemistry: Part-1 Charleton Part-227 Sodium is softer then copper but both are very good electrical canductors, (Gujranuale Board, 2008: Surgadhe Board, 2014) In solids, intermolecular forces are much stronger than liquids. Thus, it is very difficult to solids. Hence, solids have low vapour pressure than liquids. The forces in metallic solids depend upon number of valence electrons. Greater ty The forces in metalic solids depicts appear to forces. Sodium has lesser number of electrons, gil Amorphous solide like glass to also called super cooled liquid. number of valence electrons, stronger the rolless. Copper has more number of valence shell, therefore, it has weak forces. Copper has more number of valence True solids have ordered arrangement or particles, while liquids have somewhat random electrons, therefore, it has stronger forces. Thus, due to weaker forces, sodium is softer that copper Amorphous solids also have random arrangement of particles. Thus, their motecular However, both are metals and have free electrons. Hence, both are good conducting of studium is more like liquids. Hence, amorphous solids are called as super conted riquid electricity. However, due to more free electrons in Cu, it is a better conductor than ha. e.g. glass, rubber etc (4) Diamond is hard but an electrical insulator D.G. Khan Board, 2010; D.G. Khan Board, 2010; Bahasalpur Board, 2011; Cuirannels Board, 2012; Lahore Bagg 2014. P.G. Khan Board, 2007. Gujramvala Board, 2011. 2013: Lahera Board, 2009, 2013. Bahawaipur Board, 200 Sargocho Board, 2014) in diamond each carbon is sp³ hybridized. Each carbon is linked to four other carbo-Each crystal has definite arrangement of particles. Therefore a crystal can be broken amms through covalent bonds. Due to strong bonding in diamond. It is hardsally in a particular plane only, and not in any other plane However, in diamond crystal tree electrons are not present. Therefore, it cannot conduct Blus, cleavage depends upon the direction of plans. The properties, which depend electricity. Hence it is an electrical insulator. upop direction, are called anisotropic properties. Hence cleavage is an anisotropic benaviour e.g., Mica crystal consists of a large number of parallel sheets. These sheets can be easily (III) Sodium chloride and Cassium chloride have different structures. specied only parallel to their lengths. (Acad Kashmir Board, 2012) NaCl and CaCl have different structure, it is because Na* kin is smaller than Cs* kin. Thus, Na* is surrounded by only 6 Cl* ions in NaCl crystal and its co-ordination no. is only The erestale channing isomorphism mostly have the same atomic ratios 6 White due to larger size of Cs* ions, it is surrounded by B Cl. Ions. Thus, its co-ordinates number is 8. Appetailing-form does not depend upon the chamical nature of atoms. It only depends Mareover due to different sizes of Na* and Cs* ions, both NaCl and CsCl have different Pan, no of alignes, sizes of atoms and way of combination bi atoms. radius ratio Hence, substances with same no. of atoms Le. same atomic ratio, generally, have same Hence, both NaCl and CsCl have different structures. NaCl have face-pentred diffe "year structure and hence they are isomorphous. attice and CsCI have body centred cubic lattice. The translation temperature is given by elements bouling offetropic forms and compounds shouling polystorphism. (he) Judine dissolves resultly in setre chloromethane. Different allotropic forms of elements and polymorphous forms of a compound can be Lubore Source 2014) Generally poter substances dissolve in polar solvents, while non-polar dissolves in not converted into each-other by controlling temperature The temperature at which transition of one crystalline form to other organisms form polar solvents refre bigge is cuffed transpilou temberatme

one form.

(Feiselshood Boord, 2008)

compounds showing polymorphism

Since I_0 is non-polar and tetrachloromethane (CCI $_0$) is also non-polar , whice I_0 display

The capour pressure of the solids is for lass than those of liquids

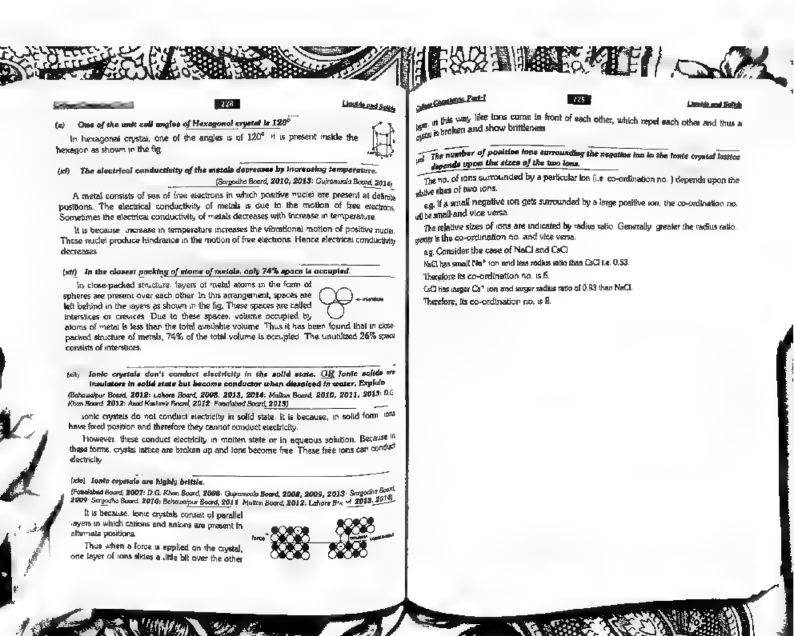
In CCL

Litterian and Solida

(Lahore Board, 2008)

Thus, concept of transition temperature cannot be used for substances, which have only

Hence, transition temperature is only given by elements showing afforcing and by







Liquide and failed

IMPORTANT DEFINITIONS

Radius Ratio

It is the ratio of the radius of cation to unloy in an ionic crystal.

i.e radius ratio = radius of cation radius of anion

It is used to express the relative sizes of different ions

Example

The radius ratio of NaCl is 0.53

The radius ratio of CsCl is 0.93

Co-Ordination Number

The number of nearest mighbours of a particle in a crystal is called its co-ordinates

e.g. In NaCl crystal, each Ne* ion is surrounded by six Cl* ions, which are present at the corners of a regular octahedron. So the co-ordination no. of Na* is six.

Similarly each CI ion is also surrounded by ski Na* ions. Its co-ordination no. is also

Co-ordination number depends upon radius ratio

Generally greater the radius ratio, greater is the co-ordination no. and vice versa.

Badots entire	Co-ordination No.
Above 0.73	8
0,414 - 0.78	8
0.22B 0.414	4
0.156 - 0.225	2

SOME CRYSTALLINE FORMS



CUBIC



CUBIC





(d) 128.5°C

(d) Metallic

HELLO! Mr Question here!

Multiple Choice Questions from PAST PAPERS

- 1 Allotropy is the property of? (D.G. Khan Board, 2010; Guyramoda board, 2011; Sang (a) compound (b) element atom (a) (d) mixture
- 2. Buy Ice (Solid CO₂) is an example of solid: (Labora bound, 1914)
 - (a) Covelent (b) Molecular (c) Ionic
 - Id. Metallic
- 3. The grystals formed due to London forces interaction are range ofic hourd, 2008, 2010) (a) lonic (b covalent ,c) molecular (d) metallic
- 6. How many allotropic forms are present in carbon (60) (a) two (b) times (c) tour (d) five
- 5 Diamond and graphite are examples of (Makes Board, 2011)
- (a) Isomorphism (b) Polymorphism (c) Isomerism (d) Allotropy
- 6. When $a = b \neq c$ and $\alpha = \beta = 90^{\circ} = \gamma = 120^{\circ}$ (Kanagandi Board, 2013) (a) Cubic (b) Trictinic (c) Hexagonal (d, Monoclinic
- 7 The Carbon atom in diamond is: Makes Boord, 2013)
 - (a) sp? hybridized (b) sp hybridized (c) sp2 hybridized (d) dsp? hybridized
- Transition temperature of tin is: (Labore Board, 2009)
- (b) 13.2°C ' (c) 0°C (a) 95.5°C
- 9 The crystal of diamond is: (Lehers Sourd, 2009)
- (b) Covalent (c) Molecular Mi ligning
- 10, Signicture of CrO₄ is (Labora Board, 2010)
- id) Triclinic (a) tetrahedral (b) octahedral (c) cubic
- 8,504 and K2CrQ4 are isomorphous solids and exist in (Sergeda-Board, Fore) [a] Table form (b) orthorhombic form (c) Trignonal form (d) Tetregonal
- 2. The transition temperature of KNOs is (Sersoche Board, 1924)
- MI 32.02°C (c: 128℃ (a) 13.2°C (b) 95.5°C
- Crystal system shown by diamond is: (Bearedper Beard, 1010)
- (д Нехадолая (b) tetragonal (c) Mosodinic (a) cobje
- 14. Attorphous solids (D.G. Ham Book, 1015)
- (8) thave sharp melting point (b) undergo cleavage when out with knife (c)
- (c) have perfect arrangement of atoms
- (d) can possesses small regions of orderly arrangement of etcms

Limite and Sulfa Charles Park 15. A crystal system in which all the axes and angles are unequal is called Gotton Board, 2007. (b) monoclinic system (a) Tetragonal system (d) cubic system (c) triclinic system 16 LiF is a crystalline substance and has (Remipted Board, 2010) (a) sonic crystals (b) metallic crystals (c) covalent crystals (d, molecular crystals 17 Which of the following is a pseudo solid? (Segodia Board, 2021) (b) NaCl (c) glass (a) CaF_e 18. Crystal of diamond is? (Supports Board, 2011) (d) metallic (b) covalent (c) molecular (n) lorde 19 Which of the following is a pseudo solid? (Sepudio Board, 2012) (b) glass (c) NaCl (d) KBr 20. Which one is not an isomorph's pair? (Falmiched Board, 2010) (a) NaNO₃ CaCO₃ (b) NaF MgO (c) K₂SO₄ K₂Ct₂O₃ (d) Zn, Cd 21 The example of hexagonal system is (Laters Seed, 2012) (a) Sulphur (b) NaCi (c) Graphile (d, Diamond 22 Transition temperature of Tin (Sn) is: (Federlobed Board, 2011) (b) 95.5℃ (c) 13,2°C (d) 32.8°C a) 128°C Answers to Multiple Choice Questions from Past Papers. Q# Ans Q# Ans Q# Ans Q# Ans Ane <u>d)</u> (b) (b)

E	16 21	,a) (c)	17 22	(c)	18	(b)	19	(b)	20	(c)
	Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in									

COLLEGE CHEMISTRY OBJECTIVE BOOK-SHORT & LONG QUESTIONS FROM PAST PAPERS

AMORPHOUS SOLIDS, CHYSTALLINE SOLIDS, UNIT CELL

Short Questions

(1) Define amorphous solid. Give example (0.0. stress Board, 2000)

(2) Differentiate between amorphous solids and crystalline solids, 150

(3) Define crystal and crystallite. (Bahmadow Bound, 2009)

Called Characters: Parts

233

15) What is a unit cell? (Perhan Board, 2006) on Draw the shape of a unit cell mentioning angles What are C.S. Khen Board, 2019 Off Describe for What are Crystallographic elements and area Crystallographic elements and area Crystallographic elements.

Define Unit cell. Draw diagram and give dimensions of hexagonal system of crystals.

(Multine Board, 2009) PROPERTIES OF CHYSTALLINE SOLIDS

Surf Questions
(1) Explain cleavage of the crystals and cleavage plane. (Bussians Boss, 2011, 2011)

Defins (a) cleavage plane (b) unit cell. (Falueband Sound, 2010)

Grophite is conductor but diamand is insulator. Give reason, grated and an

OR Why graphite is a good conductor of electricity? IS-spelle Board, 2013; OR Why graphite is anisotropic in electrical conductivity? (Receiptait Board, 2007)

Differentiate between isotropy and anisotropy). (Multip Bound, 2011)

Why do you mean by symmetry? Give elements of symmetry? (Sa

Define symmetry. Name any two symmetry elements. (Nulses Board, 2007) Write a brief note on (i) Anisotropy (II, Isomarphism, (February) 80-10)

Diffine the following terms with example. (I) Isomorphism (II) Polymorphism (II) Allotropy (iv) transition temperature (D.G. Rhan Board 2012)

Define the following terms with example. (I) Polymorphism (ii, Anisotropy (iii,

Symmetry (Gujramada Board, 2008) (10) What is habit of a crystal? (Gujramuda Board, 2909)

Define isomorphism with two examples. (Reveloted Board, 2007; Labore Board, 2007; \$1) t)

Define polymorphism with an example. Makes Board 2000

(13) Define isomorphism and polymorphism with examples (Femilies is Board, 2007; Squeened Board, 2007; Labort Board, 2013; Mallon Board, 2013) Ass. Combine SS and Side Define SS and Si

(14) Differentiate between Isomorphism and polymorphism (Nelson Search, 1918) Surg ed. 20(3)

(b) Differentiate between polymorphism and allotropy. Readpoid Board, 2012)

(17) Digits transition temperature with two examples (Resulpted Sount 2012 For 2010 D.G. Rhan Board, 2012, Malten Board, 2011, Labora Sount, 2010, 5014)

(Otto: South State Sound, 2012, Malten Sound, 2011, Labora Sound, 2010, 5014)

TORIC BOLIDS, LATTICE ENERGY

Sees Overstone

II) Why the largic crystolline solids have high melting points? (Fountehad Beard, 2002) on How the count (D.G. Khan Beard, 2009) the lonic crystalline solids have high melting points. (O.G. Klam Band, 200)

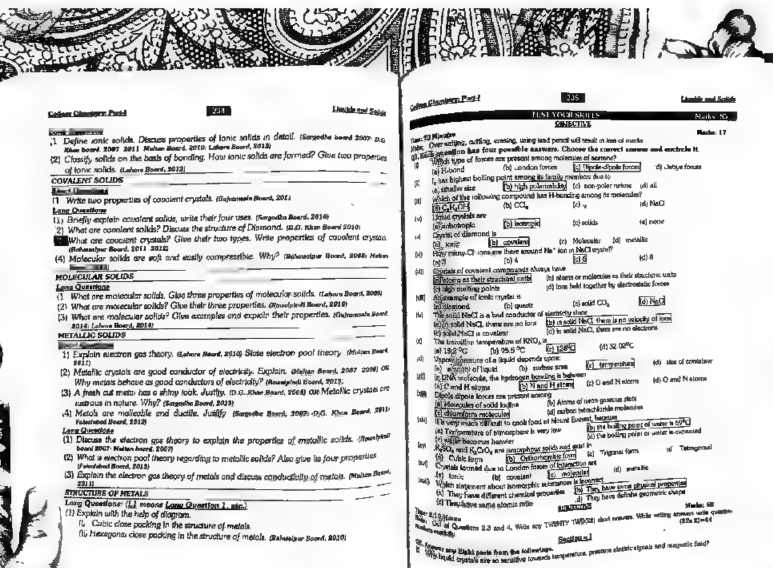
up tonic crystalline solids have high melting pouns. (AAA penn Sourd, 2018 Sour Differ Lattice energy, Give example (Labor Sourd, 2008; Mahon Board, 2018 Sour

What is Lattice and Lattice energy? (Randolm) Board, 2015

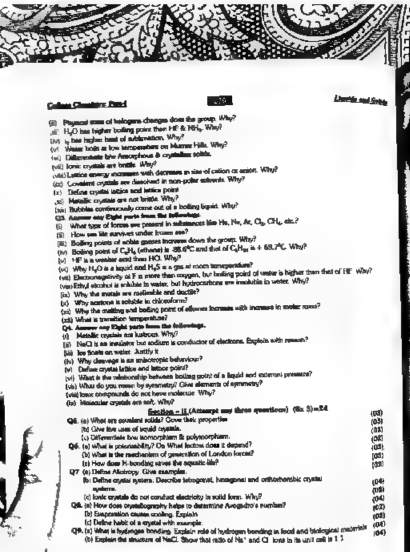
(4) Why ionic compounds do not show the phenomenon of isomerism? (Labora Board, 2010)

[5] Why ionic compounds do not show the phenomenon of isomerism? (Labora Board, 2010)

(5) What is dry toe? Mention the type of solid to which it belongs? (Rehambur Bount, \$440)
(6) NoCl is an insulator but Sodium is Conductor of electrons. Explain with remain (Regards, \$401)
Read, \$401) On Sodium is a good conductor of electricity but NoCl₁₀ is not. (Labora Board, \$401)



· 1000



hapter 5 ATOMIC STRUCTURE













Disc(CD) Surface





Atomic Structure

SUB-ATOMIC PARTICLES OF ATOM

Discovery of electron (cathode rays) as of cathode rays Discovery of proton (positive rays) Properties of positive rays Discovery of needs. Properties of neutron Measurement of change to mass ratio aim of electron Measurement of charge on electron RUTHERFORD'S MODEL OF ATOM (DISCOVERY OF NUCLEUS) PLANCK'S QUANTUM THEORY THE BOHR ATOM MODEL SPECTRUM

Continuous spectrum Line spectrum (or atom spectrum) Alonsic absorption spectrum Atomic emission apectrum Emission spectrum of hydrogen Origin of hydrogen spectrum on the basis of Defects of Sohr's atomic model X-RAYS AND ATOMIC NUMBER WAVE-PARTICLE NATURE OF MATTER (DUAL NATURE OF MATTER)

Experimental verification of dual nature of matter HEISENBERG'S UNCERTAINTY PRINCIPLE Quantum numbers Shapes of orbitals

ELECTRONIC DISTRIBUTIONS Electronic Configuration of Elements

Objective and abort answer, questions (exercise) Numerical problems (exercise)
Past Papers MCOs and Short (prestions

Test your skills

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PRATOMIC PARTICLES OF ATOM

- Metter is composed of very small particles called atom.
- According to Dalton's atomic theory, atom cannot be divided further
- However, modern researches show that atom is divisible. Several sub-atomic particles the electron, proton and neutron have been discovered

Sewery of Electron (Confinde Roys)

les Descharge Table

- It is a glass tube having two metallic electrodes sealed into it.
- it may contain a gas, air or vapours of any substance at any pressure
- It tube can be connected to a vacuum pump to maintain any low pressure.
- The electrodes are connected to a high voltage battery. The exact voltage required depends upon the length of the tube and the pressure
- A stit can be used in it to get a gharp beam of radiations.



a freed characterists for a

A neon sign' is also a discipance tube, which contain near gas at a pressure of about 10 Telegraphy is also a discipance tube, which contain near gas at a pressure of about 10 Telegraphy is also a discipance tube. for Television and computer monitor screen are also gas discharge tubes called calhode ray.
We know Ube (CRIT)

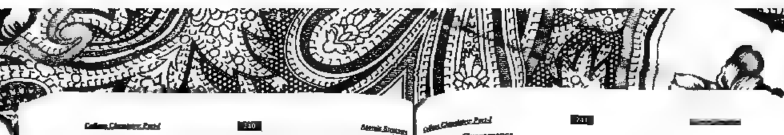
discussion of Like trans for Carlinde Ray I

Pillon Crooks Experiment

in sate 19th century, William Crooks studied the Sanose, hi used a gas discharge tube.

Standin of William Crooks Experiment

He obtained following results At normal pressure, gases do not conduct electricity even if the voltage is as high as 5000 trent. 5000 volu.



erver when the pressure inside the gas discharge tube is reduced and high voltage of 5000-10000 volts as applied, the gases begin to conduct electricity and a uniform glow appears inside the tube.

When the pressure is further reduced to 0.01 tors, uniform glow disappears and fluorescence appears on the glass walls opposite to

the cathode. This is actually due to the striking of some rays on the glass wall. These rays are called Cathode rays.

The colour of the fluorescence depends upon the composition of glass.

Different gases and vapours of different substances were used in the discharge tube. Also different metals were used as electrodes. But always same rays were produced

Properties of Cathode Ress

I Travel in a Straight Line

In 1869, Hittor' showed that Cathode says travel in a etraight .ine.

He found that cathode rays produce a sharp shedow of an opeque object placed in their path. It shows that these roys travel in a straight ling, perpendicular to the surface of cathode



2. Possess Morne

These rays can drive a small paddle wheel placed in their path. Cathode rays strike against the paddles of the paddle wheel and make it move.

This shows that cathode rays are actually begin of particles which have momentum (i.e. mass and velocity)

Entrelas Q4 (d)

The bending of the cathode rays in the electric and magnetic fields shows that they are regalitely charged.

3. Negatively Charged

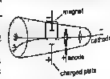
In 1895, J Perrin showed that cathode rays are negatively Charried.

He found that when cathode mys are passed through a magnetic field, these are curved downward by the magnetic field.

Muneover in 1897, J.J. Thomson showed that when cathode rays are passed through an electric field, these are deflected towards positively charged plate.

These experiments show that cathode rays are negatively charged.





They produce greenish fluorescence on striking the walls of the glass tube They also produce fluorescence in rare earlis and minerals.

e.g. <u>elumina clows red and tin stone grows vellow.</u>

5. Renduce X-rape

Cathodia revision produce X ravs, when strike an anode particularly with high atomic

6. Passes Energy (Energetic Roys)

When cathode rays strike an object, it becomes heat up showing that cathode rays are specietic rays. When cathode rays from a concave cathode fall on a platnum foll, it is in a platnum foll, it is in a platnum foll.

Innte Gosea

These can conize cases by removing electron from them. Thus, positive local are produceH

Charle Chamland Change

Triese are negatively charged. So, their addition cause reduction of a substance. Thus, these con course a chemical change.

Rive through thin foil

Desc can pass through a thin metal foil like abuninhum or gold feil

Il. Charge so moss (e/m) ratio

Their arm ratio shows that they are simply electrons.

A Thomson proved that cathods rays are actually a stream of negatively charged Periodes: He calculated their e/m ratio. He found that the e/m ratio remains same for every gas used in the discharge tube. He concluded that these are fundamental particles

Stories called these rays as electrons.

Property of Proton (Poshire Rues)

In 1896, a German physicist, <u>Fugein Coldstein discovered another type of rave valled</u>

Deliversity or canel rave, in the gas discharge tube.

ротнен

Buggin Goldstein used a periorate cathode in discharge tube. When a large voltage is applied, a low appears on the glass wall opposite to anode it is because on the glass wall opposite to the cathode tays



Triple Security of or



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Character Strategy

and after passing through perforated cathode, they produce glow on the wall.

- Since these rays pass through the holes (canals) in the cathode, therefore, these are called canal rays
- These cays are called positive rays since they carry positive charge.

Hour Canal Burs are Produced? (Origin a) Positive Roys)

These rays are produced when high-spead cathode rays (electrons) strike the the molecules present inside the gas discharge tube. <u>Cathode rays remove electrons from the gas</u> molecules and convert them into positive ions. These ions then move towards the cathode se positive ray

Properties of Positive Roys

1 Travel in a straight line

These rays travel in a straight line in a direction opposite to the cathode rays.

2. Produce Plantes (Financeacence);

They produce flashes on striking ZnS plate.

3. Postticely Charged

These rays are deflected in an electric and maintell field in a way that shows that positive charge.

4. Charge to mass (e/m) satte

- (i) The charge to mass ratio (e/m) for these raus is always smaller than for electrons.
- (ii) The elim ratio depends upon the nature of the was used in the gas discharge tabe. Heavier the gas, smaller is the 2fm value.
- (iii: The gim ratio is hishest when hydrogen is <u>present.</u> It is because, the positive partise obtained from hydrogen have least "ni" value. Hence its own ratio is highest.

Fonelistone

- rituingen produces the lightest positive particle. Rutherford called this positive particle
- as proton. It is also considered as the fundamental particle of an atom From ere ratio of proton, the mean of proton was calculated to be 1.6726 × 10⁻¹⁵ kg er 1.0073 amus.
- The mass of proton is 1836 times creater than that of electron.

Protons and Electrons were discovered until 1886 and their properties were completely independent and 1900. understood until 1895

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(d. (a): lactes Chadulck's experiment for the discovery of neutrons.

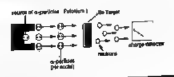
Morere of Neutron

oford Prediction

in 1920. Rutherford predicted the presence of a neutral particle neutron) in the nucleus In 1374. It is because the atomic messes of atoms could not be explained on the bads of stions and electrons only.

Chamick a Experiencent

Chadwick discovered neutron in 1932. He was awarded Nobel Prize in Physics in 1935 Discluded bombarded the nucleus of berylllura with a particles (produced from position metal source) and found that it gave highly penetrating radiation. Charged detector showed these radiations as neutral. These radiations were called neutrons



the paint frame constitution the control point of and constitution of section

Nucleaging Reactions. This neutron was produced as

Struction of a Particle from Polanium

**P6 \ He (& Particle)

Sometion of Neutron

\$80 → \$80 → BC + on (neutron)

the electrons and protons, neutrons are also considered as a fundamental particle of

September of Neutran

Comot logica gayes

Neutron san not ionise sassa bacausa it is a neutral particle

L Pres Mautran Decoya

Free neutron decays to give a proton with the emission of a neutrino and an electron

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Attent Spring

 $\sigma_{\rm s}$

Neutrons have high assestating power. They can knock out high-speed protons for paration, water paper and calculate

4. Feet and Slow Names

- Meutrons travelling with 1.2 MeV energy, are called fast neutrons
- Nutrons traveling with energy less than I eV are called slow neutrons

_, P · ae an

Sion materies are more effective than last ones for the nuclear listion process

They gard carry out marker reactions when used as projectiles. Exemples

(a) a neutron can ejects an a particle from the vocleus of nitrogen and born is produced. $\frac{1}{2}$ in $\Rightarrow \frac{36}{7}$ N $\longrightarrow \frac{13}{3}$ H + $\frac{4}{2}$ He

(ii) Story moving records produce y radiations on stilling Co. metal. In this proces sadinactive $^{66}_{26}$ Cs is convexted into $^{66}_{26}$ Zn. Neutron is captured by the nucleus of $^{66}_{26}$ Cn and $^{**}_{29}$ Co is produced. This tadioactive $^{**}_{29}$ Cu $_{\underline{emils}}$ and is attenue resolver is increased by one unit.

$$\begin{array}{cccc} \frac{1}{2} \Omega \leftarrow \frac{66}{10} \mathrm{Cu} & \longrightarrow & \frac{26}{10} \mathrm{Cu} \sim \mathrm{hv} \, \mathrm{i}_{\mathrm{T}} & \mathrm{advations}) \\ & & & & & & & & & & \\ \frac{1}{2} \mathrm{Cu} & & & & & & & & & & \\ \end{array}$$

6. Blological Activity:

These are inted in the treatment of cancer due to their buological activity

Kartlele Famous A	Cinequi(C)	Relative charge	Minus (log)	Masy fairth
Proton	+ 90Σξα B	+ .	1.5726 × 10 ⁻¹⁷	10073
Mayton	0	0	1.6750 x 10==*	1.0067
Flarron	-4.6022 x 10 ^{-m}	-1	9.,095× 10 ⁻⁴¹	5.4858 × 10 ⁻⁴



IJ Thomson determined the e/m ratio of cathode telectrons) rats in 1897. But he could act determine the charge or mass of the electrons separately.

- He subjected a beam of cathode mys in the simultaneous effects of electric and magnetic felds as shown in the figure
 - ✓ In the absence of electric & magnetic field, the electrons strike at P₁
 - When only magnetic field is applied, the electrons strike at P.
 - When only electric field is applied, the electrons strike at P.
- The strength of electric and magnetic field is so adjusted that the electrons strike at P.
- Now from the comparison of the strengths of electric and magnetic fields ofm ratio is calculated.
- The calculated value of e /m is 1.7588 \times 10¹¹ Coulomb/kg.
 It means one kg of electron carries a charge of 1.7588 \times 10¹¹ Coulombs.

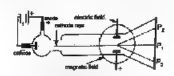


Figure - Marsina marks - 317 gledred by J. 17 canes.

Carries QS. (a):

Espiola Millikan's all drop experiment to determine the charge of an electronic

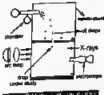
Speciment of Charge on Electron

Olifficas Oil Drop Experiment)

to 1909, R.A. Milliken determined the charge of an testion with the help of an apparatus as shown in the

Construction

- 1) consists of a metallic chamber consisting of two
- Parts, It also has two parallel electrodes in it.
- The upper electrode has a hole in it
 - The chamber is filled with air and its pressure can be





Alexande de Carlos de Carlos

adjusted with a vectoring pump

- The apparatus has an atomizer to introduce fine droplets into the chamber
- It has a microscope and are samp to see the motion of droplets.
- The falling droplet is disminated by the arc lamp perpendicular to the direction of view Thus, it appears as bright speck (spot) against dark background
- An X-ray generator is used to lonize the gas in the chamber

- Tiny droplets of all are introduced into the chamber by an element Some droplets enter into the hole
- Droplets fall under the action of gravity.
- Palling velocity of the droplets is directly proportional to its weight.

where me mass of droplet, g = acceleration due to gravity

- Now the air between the electrodes is ionised by X-rays. Oil droplets take electrons from the contred air and become negatively charged
- The electrodes are then connected to the an electric field of strength E. The oil dropists, being negatively charged, rises up towards the positively charged plates, against the force of gravity with velocity vs.

i.e
$$v_a$$
 α . Ee \sim mg _____ (2) where $e=$ charge on the droplet.

Divide eq (1) by (2)

- The strength of the electric fleid is so adjusted that the droplet becomes stand still. Interthis condition, mass of the droplet (m) can be determined.
- Thus if u_1, u_2, E, g and m are known, charge on the droplet can be determined asket equation (3)

Constusions

- Million determined charge on many of droplets and found that it was always 159 × 10⁻¹⁹ C or some multiple of it.

 This least charge 1.59 × 10⁻¹⁹ C on oil droplet is because when it picks up one electron from the air in the chamber. This value is very close to the modern value of charge which is 1.6022 × 10⁻¹⁹ C. Thus charge on one electron = 1.6022 × 10⁻¹⁹ C. Thus charge on one electron = 1.6022 × 10⁻¹⁹ C. This charge on one electron of electricity that has been determined as far.
- determined to far

Office Charthern Part !

Service QS. (c):

animie mass of electron from the above two experiments

Debraination of Moss of Electron

From e/m ratio of electrons, mass of electron can be calculated as

Since

$$\frac{e}{m} = 1.7588 \times 10^{3} \text{ C Kg}$$

$$m = \frac{e}{1.7588 \times 10^{3}}$$

$$= 1.6022 \times 10^{-19} \text{ C}$$

$$n = \frac{1.8022 \times 10^{11} \text{C kg}}{1.7588 \times 10^{11} \text{C kg}}$$

 $m = 9.2096 \times 10^{-30} \text{kg}$



Exercise Q6 (b)

http:/ord/s atomic model is based on the scattering of a-particles from a thin gold foil. Diass trans explain the conclusions.

THEREORD'S MODEL OF A TON DISCOVERY OF NUCLEUS.

- 10,1914, Lord Butherford bombarded a thin gold foll 0,00004 cm thickness with high-Speed, a particles coming from a radioactive meterial (radium or polonium).

 Absum of constricte uses obtained by clone a rightle in lead sheet. A ni
- son of a particles was obtained by using a pinhale in lead sheet. A photographic blate or a screen coated with ZnS was used as a detector. When a particles struck the street flashes of light were produced.
- Rutherford observed that most of the particles went
- though the foll.

 After were deflected at various angles and some ware deflected backward.

districtions

- Since tracet of the a-particles want straight, being therefore, most of the space in an alom is smuly.
- The central heavy region, which diffects the operations.
- hing have heavy positive charge. This part of the atom, is called Nucleus
- have heavy positive charge This part of the atom. Remaining volume of the atom is the coupled very small volume of the atom. Remaining volume of the atom is coupled by extre nuclear moving electrons.

College Chambers: Part &

27.8

Cotta cool's Atoma Mark 6

Besed on a - scattering experiment, <u>Butherford</u> proposed a planetary model of along 1911 This model was similar to solar system

- 1. An atom consists of a nucleus containing positive charge and practically the whole of its CT CESS.
- The nucleus is surrounded by a number of electrons equal to the number of protons in the nucleus.
- 3. The electrons are in constant motion around the nucleus in closed orbit like that of planets around the sun. The centrifugal force is balanced by the electrostatic force of attraction between the electrons and nucleus. Thus, electrons revolve ergund the

- 1 Sutherford's model is based upon laws of motion and travitation. These jews can be easily applied to regular bodies but not to the charged bodies such as electrons and
- 2. A revolving electron must smit engrou continuously. As a result, electron will move to a spirm path and will fall late the nucleus. Thus, whose storn would collapse. However, it never happens
- 3. If electron entits energy continuously, then a continuous spectrum



aprior disperior speak peter electronic significant less

®

PLANCES QUANTUM THEORY

in 1930. Max Planck proposed quantum theory of tight to explain emission and absorption of radiations. According to his theory, energy travels in a discontinuous maner. is composed of large number of tiny discrete (separate) units called quanta

Castellates of Placet, s On, many theory

- I. Strengy is not emitted or absorbed continuously, it is emitted or absorbed in discontinuous enganer to the form of wave packets called quanta, in see of light or ton of warmy a other called photon.
- Each wave packet or quantum has a definite amount of energy.
- A body can stoll or about among only in terms of quants.
- The energy of the quantum is directly related to the frequency of radiation by the accuration.



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Eav

of E = hv _ $_{\pm}^{(0)}$

where E = Energy of the quentum v= Frequency of radiation

h = Planck's constant=6.625 × 10⁻¹⁴ de

conclusion of Workey

k is the number of secure, which passes through a given point in one secon

into denoted by y. Its units are cycles a for all or Hertz (Pfx).

 $1\,\mathrm{Hz}=1$ cycles s $^{-1}$

to middled to wavelength as

cova or vec/a_

Where c=velocity of light=3 × 10° ms 1 1 -wavelength of any light radiation

Vice Langth

his the distance between two adjacent waste or troughs to a been of rediction

it is denote by 3.

is units are mater, nanomatre or angetrom etc.

 $1\,A^4 \approx 10^{-15}\,\mathrm{m},\,1\,\mathrm{nm} = 10^{-9}\,\mathrm{m},\,1\,\mathrm{pm} = 10^{-18}\,\mathrm{m}$

Put eq. 2 to eq. 1

Eshe/A _

 λ is related to the wave number i.e., $\tilde{\phi}_{i}$ as

λ =1 / ٧ ____

It is the number of nones per unit distance.

it is denoted by $\tilde{\mathbf{v}}$. Its units are \mathbf{w}^2 or $\mathbf{c}\mathbf{w}^3$ etc.

Put eq. 4 in eq. 3

Equations 1.3 and 5 shows that energy of light is directly proportional to its frequency

and wave number but Inversely proportional to its wavelength Greater the wave number of photons, greater is the every associated with them.

The relationships of energy, frequency wavelength, wave number about the photon of light are accepted by scientists and used by Bohr in his atomic model.

The State of the S



20%



reduce QD feets

Give the population of Bolis's charactermodel. Which postulate tells us that orbits are stationers and every a secretary

THEORY AT SOLUTIONS

in 1913. Below proposed a model of sum which removed the defects of Postsectors, see and explanations spectrum of syclenger stom. Bohr used Planck's manner ermy and purposed that electrons are present to loydrogen atom to certain quarter nery exis

- Bassion morphose-remade the maderia se distribut ration. Ends cubit has fixed energy and a
- 2. Electrical distriction and machine entering as living as a security in a fixed color. Energy is married or absorbed stop, when an electron, pumps from one orbit to exceller cities
- 8 The operator central energy whose it jumps from an outer to an inner orbit and simply smorth attention in jumps from the senser to an outer orbit.

When an disclose jumps, the energy change it Employs by

4. Elements can sensible soilly at those orbits which have fixed value of angular momentum teners. The angular researches of an eleptron in any orbit is as, singest modifice of hits.

Angeliar enomenhous — seur —
$$\frac{ab}{2a}$$

where we principal quantum number and to value is 1, 2, 2 v= reliably of electron | r = radius of the electrons

 $r={\rm sodium}$ of the electronic orbit b=F2erd/s constant = $6.625\times 10^{10}~\rm{Js}$

medically by agent, with

Thus, the parmitted values of angular moments are $\frac{h}{2\pi} \cdot \frac{2h}{2\pi} \cdot \frac{3h}{2\pi}$ etc. Hence, $\sec^{(n)}$

nt only in any one of these orbits and not in between them. i.e., angular extensions **Special**

Based on these cosmophers, Bohr presented a model of hydrogen atom and tors the H^{ℓ} -

Europe 07 Sc

Denue the equation for the redicts of 6th orbit of hydrogen atom using Bolics model

reduce of take

Consider an electron of mass 'm', and charge 'e' moving in a circular orbit of natives 'p' with velocity 'e' around the nucleus of charge Ze" The Z is the proton number (atomic number) and





of a fire change on proton.

parading to Contouts's law

gas poem by immunious managem have charged hadium in directly proportional to the dust of the magnitude of their charges, and inventely proportional to the square of succe between them.

Thus for two charges Q_{χ},Q_{χ} separated by distance (

The eclarable force of attraction between the stadeus and the electron provides the constipeted force for the motion of electron. It is given by

$$= \frac{Zex}{4\pi\epsilon_0 s^2} = \frac{Ze^2}{4\pi\epsilon_0 s^2} \qquad 0$$

where 1/4x s₀ = Contomis's into constant.

 ϵ_{0} in the vaccuum permittivity. In value is 8.84 \times 10 $^{-48}$ C 2 J 4 m $^{-1}$

The centrifugal force acting on the electron is given by

$$F = \frac{m \, v^2}{c}$$
 (2)

* For uniform decular motion, the centripetal and the conclused forms ball other Thus

$$\frac{m\varphi^2}{r} = \frac{2\pi^2}{4\pi\epsilon_0 r^2}$$

So, the equation for velocity of an electron in any odott is

$$v^2 = \frac{Z_0^2}{4\pi \epsilon_0 r m}$$
 (3)

The og shows that square of belockly of electron is inversely proportional to the radius of ontil. Il thistens electrons revolve faster in an orbit of smaller radius neuter to the nucleus. As the statement the stactron moves to higher crists of larger radius, its velocity decreases.

* According to Bohr's postulate

$$mvr = n \frac{h}{2\pi}$$

Charges: Fred

$$V = \frac{nh}{2\pi m}$$
Squarting on both rides
$$V^2 = \frac{n^2h^3}{4(1-k^2)} \qquad ... \qquad (4)$$

Comparing eq. 3 and eq. 4 , we get

$$\frac{Z_{e^2}}{4\pi c_0 m_0} = \frac{n^2 h^2}{4 \pi^2 m^2 r^2}$$

$$r = \frac{\kappa_0 n^2 h^3}{\sin Z a^2}$$

$$r = \frac{\kappa_0 h^2}{\sin a^2} = \frac{n^2}{Z}$$

$$\tau = (a_0) \times \frac{n^3}{Z}$$

iÅ≃ o™m

as =
$$\frac{s_0 \, h^2}{820 \, m^2}$$
 = constant. Its value is 5.29 x 10 $^{\circ}$ LT m, or 0.529 Å

+ Por hydrogen, Z= 1, therefore

$$r = a_{0,K} \, n^{\alpha}$$

So, the equation for radius of orbits in in an hydrogen atom is

$$r = 0.529 \times n^2$$

This eq. shows that radius of the orbit is directly proportional to the square of orbit number Hence higher orbits hape more radii and size versa. It means that radius of orbits goes on increasing with increasing orbit numbers.

→ Examples:

$$\begin{aligned} & \text{For } n=1 & r &= 0.529 \times 1^a = 0.529 \, \text{\AA} \\ & \text{For } n=2 & r &= 0.529 \times 2^a \approx 2.11 \, \text{\AA} \\ & \text{For } n=3 & r &= 0.529 \times 3^a = 4.75 \, \text{\AA} \\ & \text{For } n=4 & r &\approx 0.529 \times 9^a = 8.4 \, \text{Å} \\ & \text{For } n=5 & r &= 0.529 \times 6^a = 13.22 \, \text{Å} \end{aligned}$$

Then radius of orbits goes on increasing with increasing orbit numbers. The orbits are not equally second:

Hence, we have $-r_0-r_1 < r_0 \sim r_2 < r_1-r_0 <$

The second orbit is four times away from the nucleus than first orbit, there orbit is note The second fourth orbit is stitled times away. * (10)

Drive the formula for colculating the energy of an electron in nth orbit using Bohr's model.

Reging in view this formula explain the followings:

there of the Reculting Electron

The total energy of an electron in an orbit is equal to the sum of its K.E. due to its metion and P.E. due to electrostatic interaction with the nucleus

- Rinetto Energy

K.B. is given by

$$K.E. = \frac{2}{2} \pi n \cdot \frac{Z \, e^2}{4 \pi c_0 r \, r \alpha} \qquad \qquad \text{since} \quad v^2 = \frac{Z \, e^2}{4 \pi c_0 r \alpha} \quad \text{from: eq. (3)}$$

or K.E. =
$$\frac{Ze^2}{8\pi\epsilon_0} -$$

Potential Energy

Work done is given by

Work = Force × distance

The ejectrostatic force of attraction between the nucleus and the ejectrons is the

colourable force = $\frac{Z e^2}{4\pi \epsilon_0 \tau^2}$. If the electron moves through a small distance of the the work

done for moving electron is given by

work =
$$\frac{Ze^2}{4\pi s_0} \frac{1}{r} \times dr$$
 (work = force × distance)

In order to calculate the mate PE of elaction at a distance if from the muchaus, total work done is calculated in bringing electron from infinity to distance if Thus above eq. is thoughted hoterway the facility and the second section from infinity to distance if the above eq. is bregfaled between the limits infinity and r

$$P.E. = \int_{-\pi}^{\pi} \frac{Z_e^2}{4 \kappa \epsilon_0 r^2} dr$$

$$P = - \frac{2e^2}{4\pi\epsilon_0 r}$$



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The minus sign inclinates that the P E decreases when the electron is brought from $\mathbb{P}[\mathbb{P}]$ is sen, if this The minus sign indicates that the PE decreases unon one section is account from infinite and infinity, the electron is not attracted by any thing, thus PE is zero. At $a_{\rm E}$ At $a_{\rm E}$ is less than section $a_{\rm E}$. nearer to the nucleus, electron is attracted by nucleus, thus P.G. is less than zero

→ Thus

Total energy = E = K.E. + P.E. $E = \frac{Ze^2}{8\pi\epsilon_0 \tau} - \frac{Ze^2}{4\pi\epsilon_0 \tau}$ $\mathsf{E} = \frac{\mathsf{Z}_{\mathbf{S}^2}}{\mathsf{mt}_{\mathbf{S}^2}} \left(\frac{1}{\mathsf{B}} - \frac{\mathsf{I}}{\mathsf{4}} \right)$

$$E = \frac{Z_0^t}{\theta m t_0^\tau} \quad \underline{\hspace{1cm}} \quad (6)$$

→ On substituting the value of '>' from equation (5) in eq. (2) we gat

$$E = \frac{2a^2}{8\pi\epsilon_0} \frac{\epsilon_0 n^2 h^3}{\pi m \mathcal{E}_0^3}$$

 $\underline{E_0} n^2 h^2$ $\frac{\sqrt{m}}{\pi m Z_{\Theta}^2}$ from eq. (6)

 $E_n \simeq -\frac{8\epsilon_0^4 \, n^3 h^3}{8\epsilon_0^4 \, n^3 h^3}$ Where $E_{\rm s}$ is the energy of the electron in ath orbit.

 \rightarrow Pulting the values of $m_s *_s s_q h$

$$E = -2.18 \times 10^{-18} \times \frac{Z^2}{n^2}$$
 J

For hydrogen atom 2=1. Thus

→ The value of energy associated with 1 male of H (i.e. 1,008 g) will be

$$E_n = 2.18 \times 10^{-10} \times \frac{1}{n^2} \times \frac{6.02 \times 10^{20}}{1000}$$

or
$$E_n = -\frac{1313.31}{n^2}$$
 $i_{\text{tot}} / \text{mod}$

Hence

Charles Pand

255

$$g_{pp} = 3$$
 $E_{x} = \frac{1313.31}{3^{2}} = 145.92 \text{ ks. mod}$

For
$$n=4$$
 $E_4 = \frac{1313.31}{4^2} = -82.08 \text{ kJ} \text{ mod}$

$$\rho_{\text{off}} n = 5$$
 $\rho_{\text{dS}} = \frac{1313.31}{5^6} = -52.53 \text{ kJ} \text{ mod}$



Thus, value of energy goes on increasing towards higher orbits.

The energy differences between adjacent orbits are

$$E_{\text{B}} - E_{\text{B}} = (-145.92) + (-328.32) = 182.40 \, \text{keVmo} \, .$$

$$E_4 - 5_6 = -82.08$$
) $= -145.92$) = 63.84 ku/mol

$$\text{.e. } E_1 - E_1 > E_1 - E_4 > E_4 - E_1 >$$

The engray difference between first and infinite evergy level is given as

$$E_{\infty} - E_{0} = (0) - (-1313.31) = 1313.31 \text{ ke/mos}$$

the the contentian energy of the hydrogen. This value agrees well with the experimental

877 (1K/191

Asked Manley or dispersion of the components of while light, when it is per prior to called a spectrum.

destruit of Willia Lights

Light from sun and electric bulb consists of radiations alditerant wavelengths.

When this light is passed through a prism, it is sociated into a band of different cooper. It is because the model of loager wavelengths bend to a smaller degree while



Attack Structure

DISOUS WAR SALVO MASSELL COLUMN TO SERVE 257 255

the light of shorter wavelengths band to a greater degree and thus different colous obtained as shown in the figure

etic Spectrom:

Culture Chambers (1985)

There are seven regions in electromagnetic spectrum. One is the visible region. The spectrum of white light is visible to the naked eye & is known as <u>visible spectrum</u>. Its range is from 400 arm to 800 arm

- The rays having wavelengths below violet are Ultraviolet, X-rays and y-rays These have photons with greater energy.
- Above red are infrared, microwaves and radio waves. All these are not visible to the naked eye
- These rays form the <u>invisible spectrum</u>

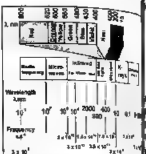


Figure | Die visit Land other zer ansiel spectrum

Types of Spectrum

There are two types of spectrum

- (I) Continuous Spectrum e.g. Rainbow
- (ii) Line Spectrum e.g. Atomic Spectrum

i. Continuous Spectrum

The operation in which rops of different wavelengths diffuse into one one into boundary can be drawn between them is called continuous spectrum.

Example, Rainbow

It is obtained from the light emitted by the oun or incandescent (electric light) solids. It is the property of znatter in bulk



What is the origin of fine spectrum?

(ii) Line Spectrum (Or Atomic Spectrum)

The spectrum in which dark or bright lines are separated by bright or dark if colled the spectrum

Origin of Line Spectrum

Line spectrum is characteristic of an atom When an element or its compo volatilized on a flame, it emits light. When this light is seen through a spectrometer, is

separated by dark spaces. The number of times and distance between them mis upon the nature of element.

Examples Example: | Une spectrum of Na consists of two yellow lines separated by a definite distance

() June 7 Agriculture of hydrogen consists of a number of lines of different colours by definite distances. In hydrogen consists of a number of lines of different colours in omnieus, definite distances. In hydrogen atom spectrum,

canos between lines decreases with decrease in wavelength and after certain wavelength, the spectrum becomes continuous.



e of Assende or Line Spectrum

nic Spectrum can be of two types.

Alamic absorption spectrum

Atomic emission spectrum

a Absorption Spectrum or Atomic Absorption Spectrum

is this, dark items are separated by bright bands.

embe: (Origin of Line [Atomic] Absorption Spectrum)

When white light is passed through a sample of a substance, it may absorb particular iors. This light on passing timough prism will form a spectrum in which dark lines will sent in place of absorbed radiations.

It has been shough in the fig.



Equal Albiet (In the unique bire)

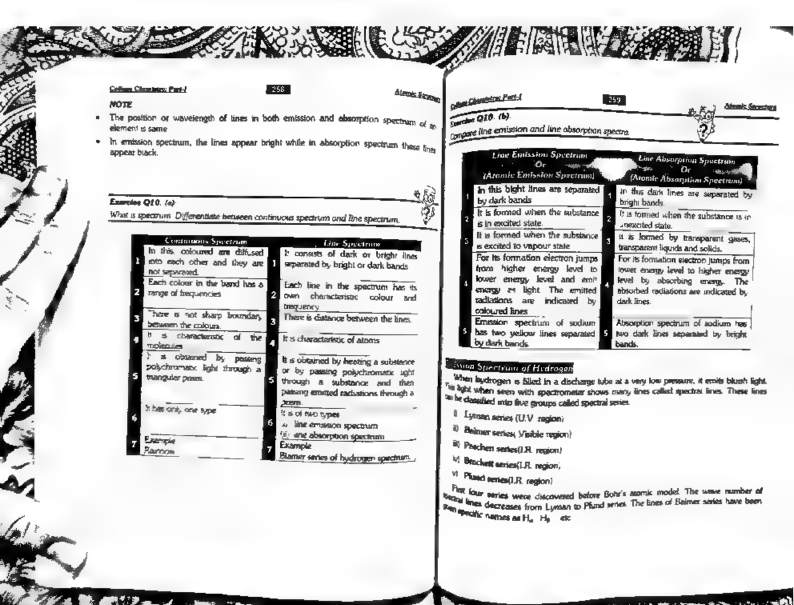
who beston Spectrum or Atomic Emission Spectrum

his, bright lines are separated by dark bands.

and IOrlain Of Atomic Emission Spectrum

When a substance is heated or subjected to electric discharge

to the radiations of definite wavelengths. These radiations will produce bright lines on a screen as shown in the figure





CHU

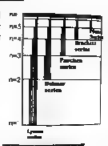
Lateral merten (ET) yearhoos	(Maria region)	(LIR region)	Henchet austen (I.R. enginet)	Etimel series (I R region)
92 per 1 30°	TOUR STOPPER	5.10× 10 ⁸	2 46 × 10°	1344 104
一种"一"	* 30 60's 10" digital.	" Bri = 10 th	3 90 × 10 ^m	R 14× 10F
2 75 - 10	23,53 c 12f \$1, text	333 × 30 ₀	4.61 × 10	<u> </u>
156 sq. 16g.	24 Ex 100 Mg hours	9.95 × 100		
130 M = 134	25.16 - 10			
100 at 1 100				

Origon Of Hedrogen Spectrum On The Besis Of Bohr's Model

According to Bohi's theory, when hydrogen atom is heated or subjected to electric discharge, its electron moves from lower orbit $\langle n_i \rangle$ to higher orbit $\langle n_k \rangle$. When this electron comes back, if ertails same energy in the form of photon of light

Five series of spectral lines are present in hydrogen atom

pecarulan.			
Series Name	a	30	Begton of Spe
Lymnon series	1	2, 3, 4,	ultraviolet
Bakmar cortes	2	3, 4, 5.	visible
Patchen series	3	4, 5, 6.	infrared
Brackett series	4	5, 6, 7	infrared
Pfunct series	5	6, 7, 8,	Infrared



Exercise Q9 (e): Derive the following equations for hydrogen atom which are related to



 $\hbar \rho$ Energy difference between two levels, n_1 and n_2

(ii) Frequency of photon emitted which an electron jumps from $n_{\rm c}$ to $n_{\rm c}$ (iii) Wave number of the photon when the electron jumps from $n_{\rm g}$ to $n_{\rm p}$

abouting of Wage Normbers of Photons of Visions Spectral Region by Bulle's Theore Larrie d'Photes.

+ According to Bohr's postulate, an electron emits energy in the form of photon when it jumps from higher energy orbit to lower energy orbit such that

$$\Delta E = hv = E_1 - E_3$$

Let energy of electron in higher energy orbit \mathbf{n}_2 is

16.5

$$E^{ij} = -\frac{8}{m}\frac{\delta_{ij}^{k}u_{ij}^{k}h_{ij}}{\delta_{ij}^{k}u_{ij}^{k}h_{ij}}$$

and, energy of electron in lower energy orbit o, is

$$E_1 = \frac{mZ^2e^4}{8\epsilon_0^2n_1^2h^2}$$

$$\Delta E = E_{2} \quad E_{1} = \frac{m Z^{2} e^{4}}{B e_{0}^{2} n_{2}^{2} h^{2}} \left(\frac{m Z^{2} e^{4}}{B e_{0}^{2} n_{2}^{2} h^{2}} \right) J$$

$$\Delta E = -\frac{mZ^{2}e^{4}}{8\epsilon_{0}^{2}n_{2}^{2}h^{2}} + \frac{mZ^{2}e^{4}}{8\epsilon_{0}^{2}n_{1}^{2}h^{2}}$$

$$\Delta E = \frac{m \, Z^2 \, e^4}{8 \, \epsilon_0^2 \, h^2} \! \left(\! \begin{array}{cc} 1 & 1 \\ n_1^2 & n_2^2 \end{array} \! \right) . \label{eq:delta-E}$$

Since for hydrogen atom Z=1 therefore

$$\text{AE} = 2.18 \times 10^{-18} J \! \left(\! \begin{array}{cc} 1 & 1 \\ n_1^2 & n_2^2 \end{array} \! \right) J$$

orginalist of Photon

→ Stace ΔE=hv

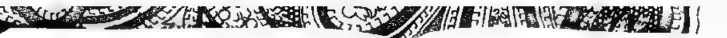
Therefore

$$\begin{split} hv &= \frac{m\chi^2\,e^4}{8\,\epsilon_0^2\,h^2} \! \left(\frac{1}{n_1^2} \! - \! \frac{1}{n_2^2} \right) \\ v &= \frac{m\chi^2\,e^4}{8\,\epsilon_0^2\,h^2} \! \left(\frac{1}{n_1^2} \! - \! \frac{1}{n_2^2} \right) Hz \end{split}$$

This sq shows that the frequency of photon emitted goes on decreasing between adjacent buts as use move towards higher orbits.

 $\frac{c}{\lambda}$ and $\lambda = \frac{1}{\nu}$, therefore $\nu = c\nu$, Hence

$$c_{V} = \frac{n_1 Z^2 g^4}{8 g_0^2 h^3} \left(\frac{1}{n_1^2} - \frac{1}{n_2^2} \right)$$



College Chamistry: Part-I

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Attento Structura

→ For hydrogen atom Z=1

$$\widetilde{\nu} = \frac{m_0 4}{8 \, e_0^2 \, c \, h^3} \! \left(\frac{1}{n_1^2} \! - \! \frac{1}{n_2^3} \right) \! = \! R \! \left(\frac{1}{n_1^2} \! - \! \frac{1}{n_2^2} \right) \! m^{-1}$$

where
$$R = \frac{m e^4}{8 \epsilon_0^2 c h^3} = 1.09678 \times 10^7 m^4 = \text{Rydberg constant}$$



Exercise Q9 (b):

Austify that Behr's equation for the wave number can explain the spectral lines of Lyman, Balmer and Paschen series

Calculations of Wore numbers Of Various Lines Present In H-Atom Spectrum

Bohr's theory can be used to calculate the wavenumbers of spectral series of emission spectrum of hydrogen atom.

Lemm Series

First line n₁=1 flower orbit, & n₂=2 (higher orbit)

$$v = 1.09678 \times 10^{7} \left(\frac{1}{n_{1}^{2}} - \frac{1}{n_{2}^{2}} \right) = 1.09678 \times 10^{7} \left(\frac{1}{1^{2}} - \frac{1}{2^{2}} \right) = 82.26 \times 10^{5} \text{ m}^{-1}$$

Second time $n_1=1$ & $n_2=3$

$$\dot{v} = 1.09678 \times 10^{7} \left(\frac{1}{n_{z}^{2}} - \frac{1}{n_{z}^{2}} \right) = 1.09678 \times 10^{7} \left(\frac{1}{i^{2}} - \frac{1}{3^{2}} \right) = 97.49 \times 10^{5} \, \text{m}^{-1}$$

Limiting line $n_t=1$ & $n_t=\infty$

$$\tilde{\nu} = 1.09678 \times 10^{7} \left(\frac{1}{n_{1}^{2}} - \frac{1}{n_{2}^{2}} \right) = 1.09678 \times 10^{7} \left(\frac{1}{1^{2}} - \frac{1}{\omega^{2}} \right) = 109.678 \times 10^{5} \, \text{m}^{-3}$$

The limiting line of Laman series lies in UV region.

<u> Edm</u>or Sorti s

First time
$$n_s=2$$
 & $n_s=3$

$$v = 1.09678 \times 10^{9} \left(\frac{1}{\eta_{1}^{2}} - \frac{1}{\eta_{2}^{2}}\right) = 1.09678 \times 10^{9} \left(\frac{1}{2^{2}} - \frac{1}{3^{2}}\right) = 25.294 \times 10^{6} \, \text{m}^{-2}$$

Second line $n_1=2$ & $n_2=4$

$$v = 1.09678 \times 10^{7} \left(\frac{1}{n_{1}^{2}} - \frac{1}{n_{2}^{2}}\right) = 1.09678 \times 10^{7} \left(\frac{1}{2^{2}} - \frac{1}{4^{2}}\right) = 20.566 \times 10^{5} \, \text{m}^{-1}$$

 $v = 1.09678 \times 10^{7} \left(\frac{1}{v_{1}^{7}} - \frac{1}{u_{2}^{2}} \right) \approx 1.09678 \times 10^{7} \left(\frac{1}{2^{2}} - \frac{1}{\omega^{2}} \right) \approx 27.421 \times 10^{6} \, \text{m}$ *The

Lighting line of Balmer senes lies in J.V. region, while other lines fall in visible region.

- Similarly wave numbers of other series of lines i.e. Paschen, Brackett and Pfund series can also be calculated.
- . Thus, Bohr's theory explained the spectrum of hydrogen atom

Exercise Q13.

Point out the defects of Bohr's model. How these defects are partially covered by dual nature of electron and Heisenberg's uncertainty principle?

Dilects of Bohr's Atomic Model

Bohr's theory explains the stability of atom, ionization energy and the spectra of tydrogen and hydrogen also atoms containing one electron e.g. He*, Li**, Be** etc. It has following defects

[] It can not explain the spectrum of multi-electron systems like He, Li, Be etc.

(ii) High resolving spectromater shows that individual lines in time spectrum of an atom actually consist of several lines, e.g. Ha-line in Ralmer series consists of five component lines. This is called ting structure or multiple lines. Bohr's theory cannot explain this fine structure. Splitting of lines shows that only principle quantum number is not sufficient. Azimuthal quantum explains the splitting of of spectral lines.

(ii) Boltz suggested circular orbits for electrons. However researches have shown that the motion of electron around the nucleus takes place in three dimensional space. Thus allowed the state of the s

atomic model is not flat

(ii) When the spectrum of atom is taken in the magnetic fletd, some new lines are created.

This is called Zeeman effect e.g. when Na spectrum is taken in a weak magnetic field, its lingle line is split up into two component lines.

straight, when emission spectrum of excited History is taken in an electrical field, lines to solid up into component lines. This is called stark effect.

Boby's theory cannot explain Zeeman affect and Starks effect.



recommendated mouthly stroom of Bests Acompe weeds t

ed in 1915 East For employer Zammer and State effect, Supermediald suppose cross species to effected white neber these in circular orbit.

in this model material is present at one of the fact of the ellipse. The elliptical push of electron goes on changing as space and thus matieus as covered by the electron cloud from all sides.



Exercise 012 (a):

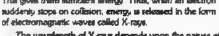
What are X-rays? What is shall origin? Flow was the idea of atomic number derived from the discovery of X-rays?

SHAYS AND ATOMIC NUMBER

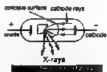
When face moving electrons strike a heavy metal anode surface in a discharge into highly courgette rays are produced. These rays are called K-rays.

erran at X co

In a gas discharge take electrons produced by sed sungeten filament are accelerated by high voltage. This gives their sufficient energy. Thus, when an electron. suddenly stops on collision, energy is released in the form



The wavelength of X-rays depends upon the nature of the target meterial Every metal has its own characteristic X-raus.



Section of Same

When X-rays are generated, they emit in all directions. These are passed through a slit and then through aluminium window. These are then thrown on a crystal of K_n [Fe[CN] μ]. which analyse the X-rays. The rays are diffracted from the crystal and a line spectrum of X rays is obtained. This is taken on a photographic plate. This X - ray spectrum is characteristic of the target material. This spectrum has discrete spectral lines. These times are grouped into K-series, L-series and M-series. Each series has various lines as K_u , K_b , L_ν L_ν M_{ac} M_p etc.

ency of a spectral line in K-ray spectrum earles as the square of atomic element acousing it. aber of the ele

$$\sqrt{v} = a(2-b)$$

This linear equation is the Moseley's Law

where v = frequency of emitted X-rays,

a = proportionality constant

Z = atomic number of element

b = screening constant of the metals



Ve for K series is plotted against 2, a straight line a obtained

This law shows that playsical and chemical properties of an electure depend upon atomic aper and not on atomic meas. Hence, Modern periodic table is based upon assess

aley's Especia

In 1913 - 1914, an English Scientist blooley studied the X-rays emissed by various He used 39 different elements from Aluminium to Gold as rarget and covered a and anyth range of 0.04 - 8 A" He obtained many enelled results

Results of Mosley's Research

- 1. The emitted rays are classified into two groups
 - (a) One with shorter wavelengths are called K series, and
 - (b) One with longer wavelengths are called L series
- 2. Wavelength of emitted X-rays decreases with increase in atomic number of target material
- The retationahlp between frequency (v) of a particular line in X-rays and atomic number of the element is given by

Importance of Moseley's Louis

- Moseley arranged K and Ar, NI and Co in a proper way in Mendeleav periodic table
- (ii) This law has led to the discovery of many elements e.g. To (43), Pm(61), Rh(45)
- (III) The atomic number of rare earths have been determined by this law

WAVE PARTICLE NATURE OF MATTER OF DEAL NATURE OF MATTER .

de Broglie Hepathesis

Einstein and Plank showed that ught has both wave-like and particle-like properties

in 1924, a French scientist, Louis de Brootle, said that all matter particles in motion also the wave-like properties. Thus, electron, proton, neutron, atoms and molecules all have both particle and wave like properties. This is called wave-parade duality

He obtained a relationship between the wavelength and the momentum of the particle. by using Einstein & Planck equations.

Einstein eq. 14

$$E = mc^4 - ... (1)$$

Plank's eq.is

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Attento Sprentino

Comparing eq. 1) and (2)

$$mc^k = hv$$

$$mc^2 = \frac{hc}{i}$$
 , Since $v=c/\lambda$

$$me = \frac{h}{\lambda}$$

$$A = \frac{h}{mc}$$

For a particle of mass (m) & velocity (v) the de Broglie equation will be

$$\lambda = \frac{h}{mv} = \frac{h}{p}$$

where p = mu = momentum of particle, h = Plank's constant' = $6.625 \times 10^{-14} \, J_{\odot}$

This equation shows that

- Particle has both wavelength $(\lambda_i$ and momentum, i.e., it has both wave and particle properties. Thus, it has dual nature.
- The wavelength of particle is inversely proportional to its momentum.

Exemples

Consider an electron moving with velocity 2.188 $\times~10^4~m~s^{-1}~$ Its wavelength can be

Mass of electron $\Rightarrow m = 9.108 \times 10^{-81} \text{kg}$

Planck's constant $= h = 6.625 \times 10^{-44} J_S$

Valocity of electron = $v = 2.188 \times 10^6 \,\mathrm{m \, s^{-1}}$

Wavelength = 1 = ?

Hence

$$\lambda = \frac{h}{mv} = \frac{6.625 \times 10^{-84}}{9.108 \times 10^{-91} \times 2.188 \times 10^6}$$

 $\lambda = 0.33 \times 10^{-9} \, \mathrm{m}$

Or
$$\lambda = 0.33 \, \mathrm{nm}$$
 , Single 1 nm = $10^{-9} \, \mathrm{m}$

- This wavelength of moving electron of first orbit of H-storn is similar to that of X-rays and can be measured.
- For a proton moving with the same vehicity, wavelength would be 1835 times smaller

Chemistry Port-I

- Similarly, art or particle moving with same velocity would have wavelength, 7344 times
- Now consider a mass of 0.001 kg (1 g) moving with a velocity of 10 m/s. Its wavelength

$$\lambda = \frac{h}{m_0} = \frac{6.625 \times 10^{-34}}{0.001 \times 10}$$

$$\lambda=6.625\times10^{-32}\,\mathrm{m}$$

This year, elength is much shorter it cannot be measured by any method

Thus, header bodies also have wavelength but their wavelength can not be measured. Thus, throld that heavier bodies do not have waves.

Esperimental Verification of Dual Nature of Matree

Dallerin und Germer Experiment

In 1927, two American scientists. Davisson and Germer gave experimental vertification of the wave nature of electron

They bombarded a thin Ni fol, with fast moving electrons. Electrons are obtained from isoted tunosten filament and accelerated by the potential difference through the charged direct. They observed that electron sidiffraction was similar to the $X \sim rays$ diffraction.

Moreover, they experimentally calculated the wavelength of electron, which was similar a that calculated theoretically using de-broglie eq.

Thus, Davisson and Germer verified de-broglie's eq. and dual nature of electron.

- Daylesson, and Germer got Nobel Prize for inventing apparatus to prove the matter
- Ds.Brogite also got separate Nobie prize for giving the equation of matter waves.

SUNHERG'S UNCERTAINTY PRINCIPLE

Il was given by Werner Helsenberg in 1927 + steles

It is impossible to determine simultaneously and procless, both position and immunity a small fast mucing particle e.g. electron

Explosection

It means the more we certain about the position of a particle, the less will be the certainty sportstend customs and vice asset

Let incertainty in position is ax and incertainty in momentum is ap then according lo Halsenbarg

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 $\Delta x.\Delta p \ge \frac{h}{4\pi}$

To satisfy this eq., it is clear that if uncertainty in position i.e., Δx is small, uncertaintymomentum i.e., ap will be large, and vice venta.

Heisenberg uncertainty principle is negligible is cone of large objects (Macroscopic objects)

Esplanation in secure of Compton Effect

This principal can be understood in terms of Compton Effect.

An electron is a very small particle. Therefore, to locate its position light having sometength shorter than its size is small (e.g. X-cays)

Heatewer, according to de-broglie equation.

i.e. Photon with abouter numericangth will featur high examentum. When such $plan_{\mathcal{K}}$ has an electron, it pushes for electron and distarbs its momentum. This is called Compa-Effect. Thus, while determining the position of electron; we will become uncertain about to more manufacture of decretor if we use light of longer was elength then determinates of he position of electron becomes impossible.

Flance we cannot determine both position & momentum of electron signaltoneously

rines with Sales's Theory

Belief in his theory assumed that electrons are material particles and revolve access to modern in some deficine orbits. Thus, their momentum and position can be determined at accuracy. But with the idea of some nature of election. Heisenberg says that it can native

Enerates Q14. (4):

Briefly discuss the some mechanical model of atom, how has it given the idea of orbid Compare orbit and orbital. (Rawalpinel Board, 2012)

ne of space to which chance of finding an electron is maximum (95%) *

Explanation: Wave Hashenford Model of Alom

- Heliamberg uncertainty, principle says that it is impossible to determine simultaneous and practicity, both position and momentum of a small fast moving particle #.9 electron
- To overcome this problem, Schrödinger, Direc and Heteroberg proposed mechanical model of atom. The best theory is that of Schrödinger

according to Schrödinger theory, electron is considered as a According to sometiments and it does not occupy a definite position in acct. Schrödinger gave a wave equation, which gives pobability of finding electrons in various regions in an atom These regions are called orbitals.

spende For H atom, the maximum probability of finding electron a as determe of 0.053 ren from the nucleus

This is the mame distance as calculated by Bohr for the 1" givit of H-astorm

de Stranders

name, for a distance shorter or longer than 0.053 nm, probability of finding electron ogates sharply in a hydrogen atom

The orbital is actually a spread of charge around the raudeus, it is often called electron

Orbit	Orbital
it is the circular path on which	It is the segion in space in which
decision revolves around the	I probability of Boding electron to
nucleus	material labout 95%,
This term is used in the Bobe's	This term is used in quantum
theory of atomic structure.	mechanical model.
lt is two dimensional	It is three dimensional
Mumber of electrons in an orbit	Each orbital contains maximum two
if given by the formula, 2n' Where 'n' is the number of orbit	electrons.
In this exact position of electron	Only probability of electron is given
is indicated.	in an orbital

ertice Q14. (6):

nd are quantum numbers? Discuss their significance

ANTEM NUMBERS

The behaviour of electrons in space around the nucleus is described by a set of four there called quantum numbers.

Quantum numbers are sets of numerical values which give acceptable solutions to indenger equation for hydrogen atom.

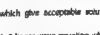
The quantum numbers are obtained by solving Schrödinger wave equation while spin which number is due to the two different orientations of electron in a magnetic field i.e., dedante & anticlockwise

Four quantum numbers and









Office Combine Page 1

1. Francisco Consultan Vinnberra

2 years micronated about Stells

- It shows the approximate distance of electron from the nucleus of an agent
- to become to the wife one n=1.23 Value of hit tells about the energy and distance of electron from the nucleus.

Consider that wakes of m. greater will be the energy and distance of electron by the ancient & vice seem. It is the measure of size of electronic shad.

- The value of a corresponds to a definite shell e.g.
- The electronism is a shell can be determined by the formula 2nd

	Shell	Copacity (2)
E	TK.	2 x 12 = 2
2	E	$2 \times 2^{r} = 8$
3	36	$2 \times 3^{\epsilon} = 13$
4	No.	2 × 4° = 32

2. Azimuthal Quantum Number ().

it gives information about Sub-shells

Origin of Asimuthal Quantum Hamber

 A spectrometer of high resolving power shows that an individual line in a line species. of an atom is actually further divided into several fine lines. It means than an individual shell is further divided into several sub-shells.

These sub-shells are explained in terms of azimuthal quantum number

- Assistable cuantum number is denoted by "\"."
- It has value from 0, 1, 2, 3 (n 1).
- The numbers 0, 1, 2, 3. corresponds for various subshells

Examples

- 0 stands for s-subshell means 'sharp'
- 1 stands for p- subshell means 'principle'
- 2 stands for d- subshell means 'diffused'
- 3 stands for f- subshell means 'fundamental'

These terms are used to describe the series of lines in the spectrum.

mber of Electrons in a Sub-Shell

it describes the shape of a subshell in which the electron is present. Maximum number of electrons in a subshell can be obtained by using the formula

the feet

 $\ell = 0$



p-subshell $2(2 \times 1 + 1) = 6$ electrons d-subshell $2(2 \times 2 + 1) = 10$ electrons

f-subshell $2(2 \times 3 + 1) = 14$ electrons

g_ige of Sab-Shells

aire of "!" also tells about the shape of orbital

A brief summany is given below

	to person secretaring a sheet I Diddie							
<u>T</u>	Substreet	Shape	Махітин приврет об Евестона					
0	5	spherical	2					
Į.	P	chambbell	6					
2	d.	\$2US/age	10					
3	£	complex	14					

the between Principal and Azimuthal Quantum No.

The relationship is as follows

	-	
o Shell	<u>t</u>	Subshell
1_K	0	s (lsl
2 L	0	s (2s)
	1	p (2p)
3 M	0	s (3s)
	1	p (3p)
-	2	d (3d)
4 N	a	s (4s,
	1	p (4p)
	2	d (4d)
	3	f_14f)etc
Dan.	4	

Schoole Omintum Numberles.

gives unformation about the different orientation of orbitals in space



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Origin of Magnetic Quantum New

- in the presence of magnetic field, lines in the line spectrum of an atom are surface spec up into verious very line lines. The appearance of these lines is explained by magnetic QUARDUM NUMBER
- It is represented by in'
- It gives the onentation and degeneracy of the orbital is space. Hence, it is also called orbital orientation quantum number
- It has value from
- m=-1 0 +1
- or m = 0, $\pm \ell$.
- The value of 'r shows the different ways in which a particular orbital can be arranged in space e.g.

Examples:

fit When $\ell = 0$. s-subshell, m=0

It shows that s-orbital can be arranged in space only in one way. Thus, it is not tuther sub-divided into other orbitals. It is a spherical and symmetrical orbital. In this <u>probability</u> a finding electron is same in all directions,

(iii When
$$\ell=1$$
 p-subshell, $m=-1,0,+1$

It means that p-orbital have three possible orientations in space. Thus, p-orbital is further sub-divided into three orbitals along X, Y and Z-axis. These are written as p., p., p., These three orbitals are present perpendicular to each other,

in the absence of magnetic field all the three p-orbitats have same energy. These are called 3-fold degenerate or triply degenerate orbitals.

(ii) When
$$\ell = 2$$
, d-subshell, $m = -2, -1, 0, +1, +2$

Thus, d-subshell is 5-fold degenerate.

(iv) When
$$\ell = 2$$
, d-subshell, $m = -3, -2, -1, 0, +1, +2, +3$

Thus, f-subshell is 7-fold degenerate etc.

Formula for number of possible orientation of orbitals

For a given value of ℓ , the total values of 'm' are $(2\ell+1)$

Thus for
$$\ell=0$$
 is subshell $(2\times 0+1)=1$ orientation

$$t=1$$
 p-subshall $(2 \times 1 + 1) = 3$ orientation

$$\ell=2$$
 d-subshelt $(2\times 2+1)=5$ orientation

$$r = 3$$
 is subshelt $(2 \times 3 + 1) = 7$ orientation



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Editionally between Principal, Azimuthal and Magnetia Quantum Nos. The relationship is given below

n.		_111		
1.	0_	_0_	18	
2	0	0	_	
-		1	2s -1 2p _z · 2p _y 2 _{D_t}	73/4
		D	2p,	Position
_		+1	20,	Crimentons of admin
3	D	0	30	
	1	-1	3p _e] गत≡
		0	3р,	Powble Crisroscions
		+1	3р,	of p-orbina.
	2	-2	3024	1
		1	3d _≠ 3d _≠	Power 1
		à	$3d_z^{-1}$	Pombir
		+ 1 ,	3d _e	Dringston
		+2	3d _{2 ,2}	el si-orbina
				_

I. Spin Quantum Numbers.

It describes the spin of election is an atom. It is denoted by s.

Olgin of Spin Quantum Number

- Alkall metals have one electron in their valence shell. When this electron jumps from the excited state to the ground state, it emits light and forms a line spectrum. High resolving performeter shows that each line in the line emission spectrum consists of two lines. This is called doublet structure. This doublet structure is different from the fine line
- structure, which is explained by azimuthal quantum number. The lines explained by azimuthal quantum number are closely
- spaced, while in doublet structure two lines are widely spaced. in 1925, Goudenit and Uhlenbeck suggested, that an electron also revolve about its exis. This is called self-rotation, it may be dockwise or anticrockwise. So an electron generates two opposite that the fields due to two opposite spins. This spin motion Produces doublet line structure in the emission spectrum of an
- alom. It can be explained by spin quantum number The spin quantum number of an electron may be + 1/2 or - 1/2



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でいか。これというから記録し

Exercises (4th treas the shapes of a. p and decripted destify these by hamping in a notes that and magnette quantum manteres

MARIE OF CHRISTALS

The shapes of orbitals can be explained on the treats of execution and respect disease unique

Consider the shapes of 5, p and d-orbitals

Sugar of a Indiana

There is only one select of energies compate number for 4 orbital. There is has only on orbital

e-contribution and a statement of the second section of the sectio

Size of a orbital increase with increase in the value of orthologic quantum number (i). Thus 2s orbital is larger in size than 1s orbital and 3s orbital is larger in size than 2s orbital.

The probability of finding electron is zero between two orbitals. This region is called nodal piene or node: surface.



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and the Cartesteen

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tempora in impace mixing these axes X, Y and Z

These three orbitals are written as p_{μ} p_{μ} and p_{μ} as shown in figure. These three orbitals e present perpendicular to each other

policial has descriptual nature. Thus it gives definite alsope to incircules. Mich profession have service shapes

See the one of a private increases with received in value of parcept administration

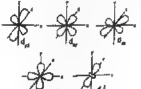


Signs of it Generales

These are from traking of comparing quantum number for d-cabital. Thus it has from mettons in space.

These five orbitals are written as d_{ny} , d_{nz} , d_{nz} , d_{nz-nl} and d_{nl} .

If the five profiteds do not have sweet shape $d_{a_i}, d_{a_i}, d_{a_i}$ and $d_{a_i - a_i}$ have four lobes each. The d_{a_i} have only two lobes and a doughthat at the centre



Shops of Forbitais is vary complicated

CANDON'S STILL

Table 5.4 Arong use u orbitals according to all

2+0=2

3+ 46

2. 3 2-3

2 . 2+1+3 0 3+0+3

4 0 4+0+4

4 . 9+1-5

4 2 4+2-6

4 3 4+3=7 5 D 5+0=3

5 2 5+2=1

5+3=6 3 : 0 6+0=6

1 6+1=7 6 2 6+2 -8

3 6-3=9

45

·Işl

ØI.

Discuss rules for the distribution of electrons in energy sub-levels and in orbitals

FLECTRONIC DISTRIBUTIONS

- 1. An orbital like s. $p_{\rm u}, p_{\rm u}, p_{\rm u}$ and $d_{\rm up}$ etc. can have maximum two electrons.
- 2. The maximum number of electrons in a shell is calculated by the formula $2n^2$ where n

There are some rules for the distribution of electrons in different sub-shells.

(D Anlino Prociple

According to Aufbau (German word, Building up) principle

Electrons are filled in autohells in order of increasing sturgy

- The energy of orbital is determined from the in+t) rule,
- Lower the (n+f) value lower will be the energy of orbital & vice VE130.
- If we orbital have same (n+t) value then lower the value of nlower will be the energy of orbital & vice versa.

Examples

Orbital	In all tours		
	in+() value	n Waiye	Remarks
44	4+0=4	4	Thus, 4s is of
3d	3+2=5	.3	
4р	4+1=5	4	lower energy
3d	3+2=5	•	Thus, 3d is of
	014-3	3	printed street of the second

Thus on the basis of this rule, order of filling of orbitals will be is 2s 2p 3s 3p 4s 3d 4p 5s 4d 5p 6s 4f 5d 6p 7s 5f 6d

2. Parties Esclusion Principle

According to this principle

it is impossible for two electrons residing in the war the same values of four quantum numbers. residing in the same orbital of a poly-electron sittle

Two electrons in the easies widted should have apposite asia $\langle \psi^{\dag}_{l} \rangle$ CXMIDE

Consider 1s orbital

For a single electron in this orbital



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ρ#1 *t*=0 m=0 & sets ⇒ + 1/2 indicated by (

Another electron enters in this orbital only if it has opposite spin to that of first. Thus for this second electron

p=1(m() m=0 & s=-1/2 andicated by 1

Thus two electrons in an orbital will have opposite spin indicated by and their one quantum mumber i.e., spin quantum number will be different

The arrangements | and | are not possible.

Orbital containing two electrons with opposite spiry is called completely filled orbital and elections are said to be paired

Orbital containing one electron is called half-filled orbital & electron is said to be espectred.

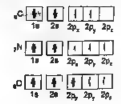
1. Inmif s Rule

According to this rule

Emblialist of some energy are available, then electrons will go in separate orbitals with ence spin, rather than in same orbital with opposite spin.

In cathier words, electrons are distributed in an atom to such a way to give maximum number of unpaired electrons.

annulés:



	tation of Elemen Table 5-5 Ltern	onte configuration of elements
Hydrogen	About number	Committee of the New York and
Helium		, 11°
Lithitem	2	182
Berytium	3	1st 2st
Boron	4	Ls2 2n2
Carbon	5	1s2 2s9 2p1 2p9 2p9
Pitrogen	6	1s 2s 2p 2p 2p 2n
	7	1st 2st 2p1, 2pt 2p3
Окуры	8	1st 2st 2pt 2pt 2pt
Fluorine	9	UF 25° 20° 200 200
Menn	10	In 2sd 2pd, 2pd, 2pd,
Sodium	11	(Ne) 3a*
tragrantum.	12	[Ne) 3e ²
Akumentee	1.3	(Ne) 3e 3p , 3p , 3p ,
Silecon	14	(Na) 34 24 24 24 24 24
Phosphorus	15	(Na) 3e ² 3p ¹ , 3p ¹ , 3p ⁴
Sulphur	16	[Ne) 3e ³ 3p ¹ , 3p ¹ , 3p ¹ ,
Chlorina	17	[Ne) 3e ² 3p ⁴ , 3p ⁴ , 3p ⁴ ,
Argon	18	[Ne] 3e ⁴ 3p ² , 3p ³ , 3p ⁴
Potamium	19	(Ar) 4x3
Celcium	20	[A] 43
Scendium	21	
iterityre	22	[Ar] 4a ² 3d ² _m 3d ² _m 3d ³ _m 3d ³ _m 3d ³ _m 3d ³ _d
Vanadium	23	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Nomite .	24	1
Longionau	25	10 m 1 m 20 m 30 m 30 m 30 m 30 m
ton	26	1 3d
chell	27	20' 30' 3d' 3d' 3d'
tectori	28	[Ar] 44 3d 3d 3d 3d 3d 3d a 3d a
opper		Pull of 20 2 20 30 30 30 30 30 .
ME		
-Burn		Ar 40 3d 3d 2d 2d 2d
Trenta		A 1 40 40 40
PARK:		1747 46° 3d1 401 401 401
Menjuni		PRI 480 State 4
Marriaday	7.0	4 do 4 do
Typion		74) # 32 4 Aur 4 7
2217	36	Ari 4a 3d 4p 4p 4p 4p

Charles Parte

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diam'r Streeter

The state of the s

OBJECTIVE AND SHORT TYSICIA, QUESTIONS (Exercise)

- of. Select the most multiple themser for the given questions.
 - The nature of positive take depends on fisher hord 20:37 (a) the nature of electrode

 - (b) the nature of the discharge tube
 - (c) the nature of the residual gas
 - (d) all of the above
- The velocity of photon is
 - (a) Independent of its wavelength
 - (b) Depends on its wavelength
 - (c) Equal to square of its amplitude
 - (d) Depends on its amplitude

idad Bourd, 2009) (D.G. Khan Board, 2011) Milan Sourd, 2011,

- (iii) The wave number of the light emitted by a certain source is 2 \times 10° m. The wavelength of this light will be (a) 500 nm (b) 500m

- (c) 200nm (d) 5 x 107 m

D.G. Khon Bearrd, 2009) (Reunalpinel board, 2011), Kiujemurale beard, 2012)

- (v) Rutherford's model of atom falled became (Nation Sound, 2013)
 - (a) the atom did not have a nucleus and electron
 - (b) It did not account for the attraction between proton and neutrons
 - (c) it did not account for the stability of the atom
 - (d) there is actually no space between the nucleus and the electrons
- (e) Bear's model of atom is contradicted by Other Best April

 - (a) Planck quantum theory (b) Pauli's exclusion principle (c) Heisenberg's uncertainty principle (d) All of the above

 - Splitting of spectral lines when atoms we subjected to strong electric field in cult-
 - (a) Zeeman effect
- (b) Stark effect
- (c) Photoelectric effect
 - (d) Compton effect
- Product Board, 2007; (Copyramoin Board, 2009; (Branspire Board, 2010; Riampired board, 2011) (Copyramoin Board, 2010) (Branspired Board, 2014) (Copyramoin Board, 2010) 2012 (Supporte Board, 2014) (In the ground state of an atom, the electron is present in the state of an atom, the electron is present.
- - (a) in the nucleus

- (c) nearest to the nucleus
- (b) in the second shell
 (d) farthest from the nucleus
- Carbonspur Board, 2009) (D.C. Khar Baurd, 1018) (Capramada bourd, 2013, 2014)
- Quantum number values for Sp orbitals are (a) p = 2, i = 1 (b) p = 1, i = 2

- (c) $\eta = 1, l = 0$ (d) $n = 2 \cdot = 0$

(a) hybrid orbitals (b) valence orbitals (c) degenerate orbitals (d) d-orbitals Februaries Boset, 2009 (Sebassipus Boset, 2009, Singarine Boset, 2009, 2012) (Ro 2019, Frankland Boset, 2011) (D.C. Khan Boset, 2012) (D.C. Khan Boset, 2012) najpindi Boord, 2009) (Madan Ba When 6d orbital to complete, the entering electron gom into (a) 7f (b) 7s (e) 7n (d) 7d bd (c) 7p (d) 7d tre Bount. 2017) (Segotha Bound, 2009) Steunbind Bound, 2010; (Gujhanouda bound, 2012) [Make May (Cujhanouda bound, 2010) FISHER AND ANSWERS TO NOT THE CHORGE OUT STIGNS Fundamental states and the secondary of game producted due to accompany of game producted due to accompany of game provides in the discontage takes in the discontage takes in the provides and the first and the secondary of the Light travels in the form of photons. Some vesocial fight is a constant quantity, therefore, velocity a photon is also constart. t-Aracter Wase larger of lagra as given by According to Partherload's allowed model, a revolving electron result and causing continuously. As a result, electron will now in a spiral path and will fall into the rescision. Thesefore, whose atom would collapse. Hence, Sutherland's model does not account for the stability of atom. $\lambda = \frac{1}{2}$ since $\nu = 2.8 \pm 0^9 \, \text{m}^{-1}$ therefore $\lambda = \frac{1}{2 \times 10^4} = 0.5 \times 10^{-6} \text{mm} = 500 \times 10^{-6} \text{m}$ =500 nm Since 1 nm =10²m (SI) Aires: (L)
When artimize spectrum of excited H-eisen is take
in an electrical field, lines are split up Into
component lines. This is called stark affect No Austria Bohr's model indicates the stact position and meanument of electron Reserver, according to Halamphery uncertainty principle both position and communitum of stactura convoid to determined wirelessed by Hence these two are apposite (contradicted) to each other. in the ground steet of an atom, the electron will be research at the downst energy level. This lowest energy level will be passent to the purious. (wiii Aus: (a) For 2p orbital, 2 stands for principle quarter number Hance for 2p orbital, n=2 According to Azisnychia quantum number, when n=2, h=0.2. Here 'O' elende for 'p-orbital' and 'I' stends for 'p' orbital, Hence, fro 2p orbital, fe 1 Thus, for 2p orbital, n=2, i=1

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animy are called

College Chambion Page 1

(ht) Orbitals having or

galant Characteristics, Part I	Z85 Acomo: So
Observing series strange are called dependence orbinals.	According to (n. 4) rule orbitals are filled energy
	wite. Thus orbitals are filled in tollowing order. Its, 2p. 3a. 3p. 4a, 3d. 4p. 5a, 4d. 5p. 6s, 44, 5d. 6p. 7s. 5f. 6d, 7p. Plance, when 6d cubital is complete. Its ordering electron will gots an 0.7p orbital.
(f. 170) in the blanks with suitable word	
ii) β-particles are nothing but mor	ving with a very high speed.
	grams.
	grams.
Epergy is when electron jumps in	from higher to a lower orbit.
i) The ionization energy of hydrogen ato	om can be calculated from mod
ağom.	
 For d sub-shell, the asimuthal quantum r 	number has value
viii) The number of electrons in a given subsi	heli si given by formula
b; The glectronic configuration of H ≤	
Answers: If electronic for 96509 (iii) 1.66's following for Bolic's (iii) 1.6	(viji) 2(2(+4)
& Indicate true or false on the case may	
A neutron is a slightly lighter particles that	m a proton
A photon is the massless bundle of energing. The unit of Rydberg constant is the recipion.	y our day momentum.
W The actual tenturis many an uther rumb	Tale and the second sec
Helecularia a a antiquata mass civile is 2000	Cable to macroscopia: volume.
the number of orbitals present in a suc-	iduced to explain accumant and make energy on of soin of electron around the nucleus.
the number of orbitals present in a such it. The magnetic quantum number was into it. Spin quantum number tells us the direction	22 v y
The magnetic quantum number was intro Spin quantum number teus us the direction	
The magnetic quantum number was into Spin quantum number teus us the direction of the state of t	学 (Iv) False (Iv) False (Iv) False (Iv) False
The magnetic quantum number was into Spin quantum number teus us the director Austress:	(v) False (v) False (v) False

or Charles Sand

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CONSTRUCTION OF THE STATE OF TH

Mr. Acap

(of Why to in necessary to decrease the presents in the discharge take to get the enthale count. 2007 Surgedia Board, 2010 Copyanicale Board, 2012)

ON Gauss do not conduct electricity at normal pressure. Why? (D.G Khan Board, 2007)

At high presence, large amount of gas is present in the discharge tube if will cause At high pressure, impressions of electrons (cathode rays). Thus, conduction of electricity is

However at low pressure amount of gas is less; therefore, electrons (cathode rays) can move and conduct electricity needly. Hence, it is necessary in decrease the pressure is

(b) Whicheour gav is seed to the discharge tube, the nature of the cathods rays remains (D.G. Khan Board, 2011 Gujranuvala Board, 2013)

Off Cathodic rays do not depend upon the nature of the gas, Discuss: (D.G. Khan Board, 2008, 2012) ON Why cathode raps are independent of nature of gas used in the discharge tube? (D.G. Khm Boss, 2009 Gueravado Bourd, 2014)

OR Why alm values of cathode mys is some for all gases: (Sargodha Board, 2007. Lahore Board, 2007.

The cathode pays are actually electrons. Since electrons are present in all atoms and their metare in some Therefore these are considered as fundamental particle of atom That notare of cathode rays remains same, no matter which gas is use. In the discharge tube.

is was proved experimentally by J.J.Thomson. He calculated e/m ratio of cathode rays by taking different gases in the discharge tabe. But he always found the same e/m ratio it shows that cathode rays obtained by different gases have some nature.

(e) Why aim value of the cothodo rope to just equal to that of electron? Foundables Board, 2007 D.G. Khan Board, 2012: Bahovolpus Board, 2010: Surgodine Board, 2018. Behavolpus Board, 2009: Culramado Board, 2012)

e/m ratio of cathodic rays is 1 7588 \times 10^{21} C/Kg, which is equal to that of electron it is became cathode rays are banically electrons

(4) The bending of the cethode rays in the electric and magnetic fields shows that the are negatively alwaysed. Solved on Page 240

(a) 19thy position rape are also autiful social rape?

(Photon Board, 2017 Asid Kashrier Board, 2012 Cultimonate Board, 2010; Bahkuralpur Board, 2018, 2010 Singulas Board, 2016- D.C. (Chan Board, 2008, Cultimosale Board, 2012)

Calculation Panel

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Engeln Goldstein used a perforated enthode for the discovery of positive rays. The Engine cathode have small pores which act as small canals. Since positive rays. The selected carrients, therefore, these are also relied canals. Since positive rays pass procedures these cameria, therefore, these are also called carral rays

g The sim orders of positive rops for different grows are different but those for cathods me the elm values to the same (Lahore Board, 2010 Rassapina Bo

on Why the properties of positive roya depend upon the nature of the gas? (Razaterial Board, 2011,

California rays are electrons. Moreover nature of electrons is same in all atoms. Thus, e/m atio is same for different gases.

Positive rays are produced due to ionization of gases present in the discharge tube e.g. consider the ionization of He and Ne

He+e" → He" + 2e" (nontine mu)

Ne+e → Ne* + 20

(positive ray)

Itahows that positive rays are nuclei of gases. Different games have different nature of nuclei. Hence, e/m ratio for different positive rays is different.

(Afte eine palue for positive rups obtained from hydrogen gas is 1836 times too the: that of cathode rays. (Lahore Board, 2007: Gujranuralo Board, 2011,

Positive rays obtained from hydrogen gas in a discharge tube consists of protons and the rathode rays consists of electrons. A proton and an electron have equal magnitude of charge bit mass of a proton is 1836 times greater than that of an electron. Hence, e/m value of Positive ray obtained from hydrogen gas is 1836 times less than that of cathode rays.

96. (e) Explain Millikan's all drap experiment to determine the charge of an electron?

Solved on Page 245

(b) What is J.J. Thomson's experiment for determining electron?

^{Solved} on Page 245

Evaluate mass of electron from the above two experiments.

Solved on Page 247

(6) (a) Discuss Chadustek's experiment for the discovery of neutrons.

Bolved on Page 243

by Rutherford's atomic model is based on the containing of a-purchase from a thin gold for

fort. Discuss it and explain the conclusions.

Selved on Page 247



Q7 (at Give the possibilities of Bole's about stockel. Which p of course in quantities? *7 -

Salved on Page 250

(b) During the equation for the radius of select of hydrogen atom uning Bole's make Saland on Page 250

(c) How does the above equation tall you that

e is allowing proportional to the aquere of the murber of crists.

(ii) Radios is immorely proportional to the number of protons in the nucleus

The equation for radius of a^a orbit of hydrogen atom, is

$$r = \frac{x_0 n^2 h^2}{m Z z^2}$$

$$cw \quad r = \frac{x_0 h^2}{m m e^2} \times \frac{n^2}{Z} \quad \text{or} \quad r \quad \alpha \quad \frac{n^2}{Z}$$

Hence, this equation shows that

- (ii) the radius is directly proportional to the square of the number of orbit (iii).

 (iii) the radius is inversely proportional to the atomic number (Z) which corresponds to the mamber of protons in the nucleus,

(d) How do you come to know that the educative of electrons in higher arbits are in then those to lower crists of hydrogen atom?

According to Bohr's theory, the equation for velocity of an electron in any orbit is

$$y^2 = \frac{Z_{\mathcal{C}^2}}{4\pi c_{\mathcal{C}^2} m} \qquad \text{or} \qquad y^3 = 2 - \frac{1}{r}$$

This equation shows that square of velocity of electron is invensely proportional in the radius of orbit (r). If means electrons revolve faster in an orbit of smaller radius nears to be nucleus. As the electron moves to higher orbits of larger radius, its velocity decreases.

(a) discribe that the distance pape between different arbits go on increasing from the loss to the higher orbits.

According to Bohr's theory, the equation for radius of orbit is

$$r = 0.529 \times n^c$$

This equation shows that radius of the orbit is directly proportional to the square of orbit number (e). Hence, higher orbits have more radii and vice verse. It means that radial of orbits goes on increasing with increasing orbit numbers.

For th = 1

 $\tau = 0.529 \times 1^{0} = 0.529 \text{ Å}$

 $r = 0.529 \times 2^{k} = 2.11 \, \text{Å}$ For n = 2

r = 0.529 x 3° = 4.75 Å For n = 3

So, we have $-r_2-r_3 \le r_3-r_3 = r_3$

Thus, radius of critits goes on increasing with increasing critit numbers.

Qd. Drive the formula for calculating the energy of m electron in site winty slobe's model. Keeping in stem this formula explain the followings:

Solved on Page 253

(a) The potential energy of the bounded electron is negative.

According to Bohr's theory. Potential Energy = P.E. = 480_aP.

The infinumation indicates that the P.E. decreases when the electron is brought from binity to distance 't'. At infinity, the electron is not attracted by any thing, thus P.E. is zero. At a point nearer to the nucleus, electron is ettracted by nucleus, thus P.E. is less than zero.

is Total energy of the bounded electron is also seguitor.

According to Bohr's theory, Total Energy = $E_n = \frac{m Z^2 g^4}{8 \epsilon_0^3 n^3 h^3}$

The minus sign indicates that the energy decreases when the electron is inought from the minus eight indicates that the energy decreases when the energy is another in another in the sample infinity to distance '?' At infinity, the electron is not attracted by any thing, thus energy is less than the point meaner to the nucleus, electron is attracted by macleus, thus energy is less than the number of the nucleus, electron is attracted by macleus. then zero i,e megative.

(i) Energy of an electron is increasing proportional to n², but energy of higher orbits are wilding greater than those of the lower orbits.

According to Bohr's theory, the energy of electron in mth orbit of hydrogen atom is given as

$$E_n = -\frac{1313.31}{n^2}$$
 kJ/mol

These, it shows

(i) The energy of an electron is inversely proportional to not

The regative sign inclinates the struction between electron and nacions. As the electron moves to higher proble, its paraction with machine decreases and hence its another increases.



Exemples

For n=1 E = - 1313.31 m / mel For n=2 E₁ = -\frac{1313.31}{2^2} = -328.32 \text{ (c) mol etc.}

Thus, the energy E_{i} is greater than E_{i}

(d) The energy difference between adjacent levels goes on decreasing sharply.

mZ^ze⁴ According to Bole's theory, Total Energy = E_n =: $8s_0^2n^2h^2$

This equation shows that the energy of an electron is inversely proportional to π^{α} H regy difference between adjacent levels goes on decreasing sharply

The energy differences between adjacent orbits are

 E_4 $E_6 = (-328.32)$ [-1313.31] = 984.99 kl/mol $E_0 - E_4 = (-145.92) + 328.32) = 182.40 \text{ kH/mad}$ $E_4 - E_9 = (-82.08) - (-145.92) = 63.84 \text{ kJ/mol}$

Thus, $E_2 - E_1 > E_1 - E_2 > E_4 - E_5 >$

Q7. (a) Durker the following equations for hydrogen stom which are related to: & Energy difference between two levels, n, and n_f.

(iii) Frequency of photon emitted which an electron jumps from n_k to n_1

(iii) Wase number of the photon when the electron funces from n_2 to n_1

Saland on Page 260

(b) June(f) that Bohr's equation for the same number can explain the specied last famous, Solmer and Panches series.

Solund on Page 262

Q10. (a) What is spectrum. Differentiate between continuous spectrum

Solved on Page 258

(b) Compare the emterior and the absorption spectra

Solved on Page 259

(a) What is the origin of line spectrum?

Solved on Page 256

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(1) (d) Hydrogen atom and He* are monoelectronic system, but the size of He* to much smaller than H. Why?

Both hydrogen atom and He* ion have one electron in their outermost shell. However, Both hydrogen again and the first have one spectron in their outermost shell. However, the raideus of He* has greater positive change(due to two protons) than that of hydrogen atm (due to one proton). Therefore, nucleus of He* attracts its electron more powerfully as compared to hydrogen. Hence, size of He* becomes smaller than hydrogen.

(b) Do you think that the size of LI* is oven smaller than No? Justify with calculations

Both He" and Li^{p*} sons have one election in their outermost shell. However, the nucleus of Li' has greater positive charge(due to three protons) than that of He' son ,due to two roton). Therefore, nucleus of Li²⁺ attracts its electron more powerfully as compared to He' hence, size of Li²⁺ becomes smaller than hydrogen.

Nathernatical Calculations

For monoelectronic systems, Bohn's equation for radius of arbit is given by

$$t = \frac{n^2}{2} \times a_0$$

Where

Z = Atomic number & $\eta = \eta combes of orbit$ a_s= constant = 0.529 Ast Thus for 1st orbit of 15st

Thus for
$$1^n$$
 orbit of 1^{n-1}
 $n \Rightarrow 1$ $Z = 3$

$$r = \frac{1^2}{3} \cdot 0.529 = \frac{0.176 \text{ Å}}{2}$$

For Jª orbit of He' ion

$$r = \frac{1^3}{9} \times 0.529 = 0.264 \text{ Å}$$

It shows that size of L ** is smaller than He* Q12. [a] What are X-ray of What is their origin? Most ones the idea of or

derived from the discovery of K raye?

Salved on Page 264

h) Hou does the Bohr's model purity the Moreley's equation?

According to Bohn's theory, the frequency of emitted photon is given as

$$v = \frac{mZ^3e^{4/3}}{B_{coh}^3} \left(\frac{1}{m^3} - \frac{1}{m^3} \right)$$



or vαZ[‡]

o Frez

Thus, when electron jumps from higher orbit 'ng' to lower orbit 'tra', the square root of elements of bytomor emisses as givently bendouspoury to the securic settinges of the ejement och is the Mountay's law i.e.,

w Jaz

Q23. Point out the defects of Sale's model. How these defects eve p dest nature of electron and Heleschung's uncertainty principle?

Salmed on Page 863

Q14. m) Bright discuse the wave association model of otom. Note has it gives the lit-of orbital? Compare white and orbital. (Resolpted Board 202)

Solved on Page 268

manhesi? Discuss that styrifts (b) When my quant

Saloed on Page 209

hal quantum number, 5 k 3. Sergocho floord, 2019

The values of magnetic quantum number (m) is related to asknothal quantum number (6) by the relation

Hence

When 6 = 3 k is f-orbital. & m = 3, 2, 1,0,+1,+2,+3

Thus parties 7 values for $\ell=3$

These seven values show that f-orbital has seven different orientations in spo

Q15 (a) Discour rules for the distribution of electrons in energy sub-levels and b

Soloed on Page 276

(b) What is (a + § rule. Arrange the orbitols according to this rule. Do you think the title rule is applicable to degenerate orbitols?

The orbitals of a subshell having some (n+ f) value are called degenerate orbital country have some they have some because they have same energy, Hence, (n+ f) value are called degenerate orbitals. However, decements orbitals have been politically because the applied to degenerate orbitals. orbitals. However, degenerate orbitals have different values of magnetic quantities

Charles Park

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Alberts Street,

therefore, they can be differentiated by m values, e.g. 2p., 2p., 2p., are published orbitals. Their (n+4 are seme i.e., (2+1)=3 but their in values are different incomes orbitals.

Distributor characters in orbitals of sales, sales, sales, sales, sales, sales

ala 3et 2pt 3pt 3pt 4et 3dt 4pt 5et 4dm 5pt 6et 40 5dt

TO THE ROLL OF THE RESTREET

Ca 16" 28" 20" 38" 30" 4a" 36 "

And 12 22 20 32 30 42 300 60 50 400 50 60 40 500

MCr 1st 2st 2pt 3st 3pt 4st 3dt

15" 25" 2pt 3el 3pt 4el 3din 4pt 5el 4din 5pt

"Ba 15" 25" 25" 35" 36" 46" 34" 40" 55" 46" 50" 65" 41" 56" 65"

\$16. Dream the shapes of a, p and d-orbitals. Justify these by imaging in about the ministral and magnetic quantum numbers

Salved on Page 274

1 IMPORTANT FORMULAS berg (5), frequency (4), wavelength (1), $Z e^2$

transfer (* June mass interesperations)

 $l = h_{k} - c_{m,k}$ k = 1, $k = mc^{2}$

da's confloor.

paya momentum = mv. ∈ 2×

क्षेत्र वर्ष का क्रकी

 $= \frac{2\pi h^2}{4m_0 e^2} \times \frac{n^2}{2} Or$

<u> विकास को का क्षेत्रकार</u>

Or $\hat{\mathbf{E}} = -2.8 \times 0^{-11} \times \frac{2^{-1}}{2}$

Dr 2 _ 331

beautiber of a photon

 $\frac{1}{16\frac{2}{3}6\frac{1}{6}} \sqrt{\frac{1}{n_{1}^{2}}} \cdot \frac{1}{n_{2}^{2}} = \frac{1}{n_{2}^{2}} + 2\left(\frac{1}{n_{1}^{2}} + \frac{1}{n_{2}^{2}}\right)$ bg PR - 75'

 $K = \frac{8\pi r^2 L}{\kappa^2 e}$

Months a popular N= 6(2 b)

 $\lambda = \frac{h}{mv} = \frac{h}{p}$

Permitted product to the control of Chans on an electron e=1.6022 × 10 C

Mass of an electron m = 9 109 × 10 11 kg Velocity of Bohl c =3 × 10° ms

Rudberg constant

 $R = \frac{m_e^4}{8 g_e^4 c_h^4} = 1.09678 \times 10^7 cs^4$

and a₀=0.529 Å

中心のなるので、アージャンプの一般





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0

NUMERICAL PROBLEMS (Exercise)

(a) A photon of light with energy 10⁻¹⁹ J is emissed by a source of light (a) C this energy two the assertingth, frequency and some number of the when

$$\begin{array}{l} F = 10^{-16} \ J \\ h = 6.625 \times 10^{-34} \ Js \\ c = 3 \times 10^6 \ m/s \\ v = ^4 \\ A = ^7 \end{array}$$

Or
$$v = \frac{E}{h} = \frac{10^{-19}}{6.625 \times 10^{-34}} = \frac{1.509 \times 10^{16} e^{-1}}{1.509 \times 10^{16} e^{-1}}$$

Since
$$\lambda = \frac{c}{v} = \frac{3 \times 10^{4}}{1.509 \times 10^{14}} = 1.988 \times 10^{16} \text{ m}$$

Since
$$v = \frac{1}{\lambda} = \frac{1}{1.988 \times 10^6} = \frac{3.080 \times 10^6 \text{m}^{-1}}{1.000 \times 10^6 \text{m}^{-1}}$$

(b) Convert this energy of the photon into ergs and calculate the wave length is a frequency in Hz and cover number in \cos^{-1} in $a = 6.625 \times 10^{-24} \, \mathrm{Js}$ $c = 3 \times 10^{4} \, \mathrm{m/s}$ $c = 10^{-11} \, \mathrm{Js} + 10^{24} \times 10^{7} \, \mathrm{erg}$

$$h = 6.625 \times 10^{-14} \text{ Ju}$$
 $c = 3 \times 10^{4} \text{ m/s}$
 $E = 10^{-19} \text{ J} = 10^{31} \times 10^{7} = 10^{-19} \text{ and}$

$$\varepsilon = 3 \times 10^6 \ \text{m/s} = 3 \times 10^{16} \ \text{cm/s}$$
 , $\nu = 7$

$$\lambda = 7$$

$$\nu = \frac{E}{b} = \frac{10^{-18}}{6.625 \times 10^{-29}} = \boxed{1.509 \times 10^{16} e^{-1}}$$

$$\lambda = \frac{c}{v} = \frac{3 \times 10^{30}}{1509 \times 10^{31}} = \boxed{\text{L.988} \times 10^{-6} \text{ cm}}$$

$$v = \frac{1}{\lambda} = \frac{1}{1.968 \times 10^{-4}} = \frac{8.080 \times 10^{9} \text{cm}^{-1}}{1.968 \times 10^{-4}}$$

QIA. The forwards Jos

$$E_a = \frac{m_B^4}{8\epsilon \delta n^2 h^2}$$

Calculate the many of electron in first orbit of hydrogen

Salations

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Ç

$$\begin{split} z_0 &= 8.85 \times 10^{-22} \, \text{C}^{-1} \, \text{J}^{-1} \, \text{m}^{-1} \\ h &= 6.625 \times 10^{-24} \, \text{Js} \\ m &= 9.1 \times 10^{-21} \, \text{lrg} \\ e &= 1.6022 \times 10^{-19} \, \text{C} \\ E_1^{\text{sec}} ? \\ 0 &= 1 \end{split}$$

Energy in the first orbit is given by

$$E_1 = -\frac{m_2^4}{8 s_0^2 n_0^2 h^2}$$

$$E_1 = \frac{(9.1 \times 10^{-31})(1.602 \times 10^{-19})^4}{8 \times (8.85 \times 10^{-12})^4 \times (6.625 \times 10^{-34})^4}$$

$$E_1 = 2.18 \times 10^{-11}$$

Q19. Bahr's equation for the radius of ath orbit of electron in hydrogen stom to

$$r_n = \frac{\varepsilon_0 n^2 h^2}{\sin \varepsilon^2}$$

Walls doing calculations take ours of units of source parameter

(i) When the electron moves from n=1 to n=2, how much does the radius of the orbit

Radius of nth orbit is given by

$$L_{\mu} = \frac{\sin \phi_{\frac{1}{2}}}{\sin \mu_{\frac{1}{2}}} \times \mu_{\frac{1}{2}}$$

$$\mathbf{v}_{n} = \frac{\left(8.85 \times _{\star} 0^{-12} C^{2} \phi^{-1} m^{-1}\right) \left(6.625 \times 10^{-19} d_{\star}\right)^{2}}{9.14 \times \left(9.1 \times 10^{-91} M_{\odot}\right) \left(1.602 \times 10^{-19} C\right)^{2}} \times n^{2}$$



 $a_0 = 0.529 \times 10^{-10} \text{m} \times n^2 = 0.529 A^4 \times n^2$

Thus the work 4 = 0.529 × 19 = 0.529 Å

Thes for n= 2

Phone transports value = $t_0 - t_1 = 2.11 \text{ Å} - 0.519 \text{ Å} = [3.51] \text{ Å}$

Smor & = 0.529 × 10: "m × n³ = 0.529 Å × n³

10 mm = 1 Å

 $\eta_1 = 0.529 \times 2^2 = 2.11 \text{ Å}$

Thus for n=3

h = 0.529 × 3F = 4.75 Å

Hance deserte territoi=q-q-4.75 A* 2.11 A*= 2.66 A

Also for a=9

1, = 0.579 · F = 42.869 Å

And tor n= 10

rm = 0.529 × 10° = 52.9 Å

Printer democra travelled = 1₁₁ 2,=\$2.9 Å 42.849 Å ≈10.05 Å

Q28 Amount for following quantum, by performing the calculation



Epongs of electron as rith crisis in given by

Where $\overline{\epsilon}$ = atomic number -n = number of orbit

Collabora Aust 1-1

$$E_{\rm p} = (3.18 \times 10^{-10} \times \frac{1}{\rm m^2})$$

Thus

Form = 1 | Er = #16 \ 10 4 \ 1 2 4 16 \ 10 41

 $For n = 2 \cdot E_{\rm f} = 2 \cdot 18 \cdot 10^{-14} \cdot \frac{1}{2} = 5.45 \cdot 10^{-12} - 0.545 \cdot 10^{-14}$

For n = 3, $E_1 = -0.16 \times 10^{-10} \times \frac{1}{45} = -0.43 \times 10^{-10} 2 \times -0.201 \times 10^{-10} 3$

For n = 4 Eq. = $2.18 \times 10^{10} \times \frac{1}{12} = -1.36 \times 10^{10} J = -0.36 \times 10^{10} J$

For n = 5 $E_5 = -2.18 \times 10^{-16} \times \frac{1}{5^4} = -8.72 \times 10^{-16} J = -0.0872 \times 10^{-16} J$

Energy differences will be

 $E_{e} = E_{g} = (-0.545 \times 10^{-10}) = (-3.18 \times 10^{-10}) = 1.636 \times 10^{-10} \text{ J}$

$$\begin{split} & - \epsilon_0 = -0.365 \times 10^{-14} + (-0.545 \times 10^{-14} = 0.303 \times 10^{-16} J \\ & E_0 = E_0 = (-0.136 \times 10^{-14}) + (-0.545 \times 10^{-16} = 0.303 \times 10^{-16} J \\ & E_0 = (-0.136 \times 10^{-14}) + (-0.242 \times 10^{-16}) = 0.0488 \times 10^{-16} J \\ & E_0 = E_0 = (-0.0872 \times 10^{-16}) + (-0.136 \times 10^{-16}) = 0.0488 \times 10^{-16} J \end{split}$$

(I) that energy differency behavior second and shird orbits to approximate that the behavior first and accord orbits.

Energy difference between E_a - E_b and E_e - E_b . The ratio of the anergy difference between E_a - E_a and E_a - E_a is given by

 $\frac{E_2-E_1}{E_3-E_2} = \frac{1.635 \times 10^{-16}}{3.03 \times 10^{-16}} = 5$

or 1 (E2 - E3) = (E3 - E8)

Hence, energy difference between $E_d = E_0$ is app than Eg Es

(c) Calculate the energy of electron in He' in first flow orbits and fuelify that the a afference are different from those of hydrogen atom

$$\underline{p}_{\rm d} = -2.18 \times 10^{-19} \times \frac{Z^{8}}{{\rm g}^{1}} = -2.18 \times 10^{-19} \times \frac{2^{3}}{{\rm g}^{1}}$$

For
$$n = 1$$
 $E_1 = -2.18 \times 10^{-10} \times \frac{2^4}{1^2} = -6.72 \times 10^{-10} d$

For
$$n=3$$
 $E_0=-2.16\times 10^{-10}\times \frac{2^6}{3^7}=-9.66\times 10^{-10}\,\mathrm{d}=-0.968\times 10^{-10}\,\mathrm{d}$

For
$$n=6$$
 E₄ = $-2.16 \times 10^{-10} \times \frac{2^{9}}{6^{9}} = -3.488 \times 10^{-19} J = -0.3488 \times 10^{-10} J$

$$E_{\rm s} = E_{\rm s} = .0966 \times 10^{-41} = (.218 \times 10^{-43}) = 1.21 \times 10^{-43}$$

Energy differences will be $\begin{array}{lll} E_{a} & E_{b} = -0.2 \ 18 \times 10^{-10} \) & = -0.872 \times 10^{-10} \) & = -0.54 \times 10^{-10} \ 4 \\ E_{a} & E_{b} = -0.966 \times 10^{-10} \) & = -0.218 \times 10^{-10} \) & = 1.21 \times 10^{-10} \ J \\ E_{a} & E_{a} = -0.546 \times 10^{-10} \) & = -0.966 \times 10^{-10} \) & = 0.423 \times 10^{-10} \ J \\ E_{a} & E_{a} = 1.0.3486 \times 10^{-10} \) & = -0.546 \times 10^{-10} \) & = 0.196 \times 10^{-10} \ J \\ J denote difference of energy between the energy inveit of 14e^+ is different into$ hydrogen

(d) the year which that proups of the spectral times of the' are as different plants that those for hydrogen about give reasons

Since amongs differences between energy levels in 11c° ion is different for Tuyloogen, therefore spectral from in 11c° ions spectrum will be different from fact of tuylorum. ANSWERS SHARE FARMS

(8) Colordon the color of principal quantum marker if an electric in hydrogen produce to an orbit of energy - 6.242 - 20 rd S.



Energy of electron in oth orbit is given by
$$g_{\rm el} = -2.18 \times 10^{148} \times \frac{2^3}{\kappa^2}$$

Hence
$$0.242 \times 10^{-18} = -2.18 \times 10^{-19} \times \frac{1^2}{n^2}$$

$$n^2 \approx \frac{-2.16 \times 10^{-18}}{-0.242 \times 10^{-18}} \approx 9$$

022. Bohr's formula for the energy lacels of hydrogen stom for any system say 11, 142°, 13 2° ato is

$$E_{n} = \frac{m Z^{2} e^{4}}{8 \epsilon_{0}^{2} n^{2} h^{2}}$$

$$E_n = K\left(\frac{Z^2}{n^2}\right)$$

For topdrogen Z=1 and for He^{*} , Z=2.

(4) Draw an energy lovel diagons for hydrogen nion and He'.
(b) Tituliting that S = 2.8 × 10⁻¹³ d_c calculate the energy near electron from hydrogen atom and from He'.

$$K=2.18\times10^{-10}\,\mathrm{s}$$

For toydrogen Z = 1

Emergy in ruth orbit is given by

Form = 1 Ear 2 18 = 10 14 (1) 2 2.18 - 10 14 4

For
$$n = \infty$$
 $= \frac{2}{8} \cdot 10^{-10} \left[\frac{I^2}{\pi^2} \right] \circ 84$

Marica: to move an election from hydrogen's time what to all infinite distance, the

E. E. = 0 - (-2.18 - 10 -) = 2 16 × 19 19

Thre is the ionization energy of hydrogen atom



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9

Por He* ion Z = 2 Energy in 18th arbit is given by

$$\underline{\mathbf{g}}_n = -\mathbf{K} \left(\frac{\underline{\mathbf{Z}}^k}{n^T} \right)$$

 $E_{\rm e} \simeq -2.18 \times 10^{-16} \bigg(\frac{2^9}{1^6}\bigg) = -8.72 \times 10^{-18} \, J$

Por
$$n = \infty$$
 $E_n = -2.18 \times 10^{-18} \left(\frac{2^3}{m^2}\right) = 0.0$

Hence to make an electron from bits' lon's first orbit to an infinite distance, the energy required will be

$$E_a - E_1 = 0 - (-8.72 \times 10^{-10}) = 8.72 \times 10^{-103}$$

This is the ionization energy of He* ion.

(a) How do you furtify that the energies calculated in (b) are the tanization energies of it and He'?

The amount of energy required to remove an electron from an atom or in

to an infinite distance is called ionization energy. Hence 2.18 \times 10 $^{-18}$ J and 8.72 \times 10 $^{-18}$ J are the ionization energies of Hence 2.18 \times 10 $^{-18}$ J are the ionization energies of He atom and He* (on respectively-

(d) Use Avogadro's number to convert tonication energy values in kilmot' or H and He

The ionization energy of H-atom in ki/mol is given as

E =
$$2.18 \times 10^{-18} \times \frac{6.02 \times 10^{28}}{1000} = \frac{1312.36 \text{ kJ/mol}}{1000}$$

The ionization energy of He*-ion in kJ/mol is given as

$$E = 8.72 \times 10^{-18} \times \frac{6.02 \times 10^{23}}{1000} \times \frac{5249.4 \text{ kJ/mol}}{1000}$$

(a) The experimental solute of lontgetton energy of it and He* are 1331 tol mol¹ of 6250 kJ mol³ respectively. How do you compare your values with experimental values?

The calculated values of ionization energies for H-atom and He^{*}-ion using Bohr's theory are 1312.36 kJ/moi and 5249 4 kJ/moi respectively.

These results agree well with the experimental model in 1931 kJ/moi for h

These results agree well with the experimental results i.e. 1331 kilmol for its mand 5250 without factions. atom and 5250 ml/mot for He*ton.

Both H-storn and Harton consists of one electron each and the above reals clearly shows that Bohr's theory is perfectly applicable to one electron system.

Acres Parts

083. Colculate the trave number of the photon others the electron jumps jo (1) n = 5 to n = 2.

(fi) n = 5 to n = 1.

In which earles of spectral lines these photoes will app

(aletton)

Rydberg constant = $R = 1.097 \times 10^7 \, \text{m}^{-1}$

When electron jumps from $n \neq 5$ to n = 2.

The wave number of the photon is given by the eq.

$$\widehat{c}_{i} = B \left(\frac{1}{n_{1}^{2}} - \frac{1}{n_{2}^{2}} \right) = 1.097 \times 10^{7} \left(\frac{1}{2^{2}} - \frac{1}{5^{2}} \right)$$

$$5 \approx 1.097 \times 10^7 \left(\frac{1}{4} - \frac{1}{25}\right)$$

$$J = 1.097 \times 10^7 \times \frac{21}{100} = 2.30 \times 10^4 \, \text{m}^{-1}$$

This spectral line is present in visible region(Balmer Series) When electron jumps from n = 5 to n = 1

The wave number of the photon is given by the eq.

$$c = R\left(\frac{1}{n_1^2} - \frac{1}{n_2^2}\right) = 1.097 \times 10^7 \left(\frac{1}{16} - \frac{1}{5^2}\right).$$

$$\bar{\nu} = 1.097 \times 10^7 \left(\frac{1}{1} - \frac{1}{25}\right)$$

$$\bar{c} = 1.097 \times 10^7 \times \frac{24}{25} = 1.06 \times 10^7 \, \text{m}^{3}$$

This spectral line is present in UV region (Lymen Series)

Q24: A photon of a more number $102.70 \times 10^9 \, \mathrm{m}^{-1}$ jumps from higher to n=1Salation.

Rydberg constant = $R = 1.097 \times 10^7 \,\text{m}^{-1}$ $\frac{n_1 = 1}{n_2 = 2}$

 $\alpha_s =$ The wave number of the photon is given by the eq.

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Alonic Street

$$v = R\left(\frac{1}{n_1^2} - \frac{1}{n_2^2}\right)$$

$$102.7 \times 10^3 = 1.097 \times 10^7 \times \left(\frac{1}{1^2} - \frac{1}{n_2^2}\right)$$

$$\frac{102.7 \times 10^5}{1.097 \times 10^7} = 1 - \frac{1}{n_2^2}$$

$$0.93637 = 1 - \frac{1}{n_2^2}$$

$$\frac{1}{n_2^2} = 1 - 0.93637 = 0.0636$$

$$n_2 = \frac{1}{0.0636}$$

$$n_2 = \sqrt{\frac{1}{0.0636}} = \frac{3.996 \times 4}{1}$$

(b) Indicate the name of the series to which this photon belongs.

This spectral line is present in Lyman series

(c) If the electron will fall from higher drift to n = 2, then valculate the wave number of the photon emitted. Why this evergy difference is so small as compare to above calculations?

When electron jumps from n = 4 to n = 2.

The wave number of the photon is given by the eq.

$$v \approx R \left(\frac{1}{n_1^2} - \frac{1}{n_2^2} \right) \approx 1.097 \times 10^7 \left(\frac{1}{2^2} - \frac{1}{4^2} \right)$$

$$v \approx 1.097 \times 10^7 \left(\frac{1}{4} - \frac{1}{16} \right)$$

$$v \approx 1.097 \times 10^7 \times \frac{3}{16} = \frac{2.056446 \times 10^6 \text{ m}^{-1}}{16}$$

Energy difference for n=4 to n=1 can be calculated by the eq

$$\Delta E = 2.18 \times 10^{-14} \left(\frac{1}{r_0^2} - \frac{1}{r_0^2} \right)$$

Thus
$$\Delta E = 2.18 \times 10^{-18} \left(\frac{1}{1^2} \times \frac{1}{4^2} \right)$$

$$\Delta E = 2.18 \times 10^{-18} \left(\frac{1}{1} - \frac{1}{16} \right)$$

$$\Delta E = 2.18 \times 10^{-18} \times \frac{15}{16} \times \frac{2.04 \times 10^{-18}}{16}$$

And the energy difference for n=4 to n=2 can be calculated by the eq.

Thus
$$\Delta E = 2.18 \times 10^{-18} \left(\frac{1}{2^2} - \frac{1}{4^2} \right)$$

 $\Delta E = 2.18 \times 10^{-18} \left(\frac{1}{4} - \frac{1}{16} \right)$
 $\Delta E = 2.18 \times 10^{-18} \times \frac{3}{16} = 4 \times 10^{-19} \times \frac{0.4 \times 10^{-18}}{10^{-18}}$

The energy difference in second case is small,

It is because electron travel more distance from n=4 to n=1 than n=4 to n=2. And since energy is directly related to the distance of the electron, hence energy difference in second case is smaller than first case.

Q25. (a)What is de Broglie's wavelength of an electron travelling at half a speed of light?

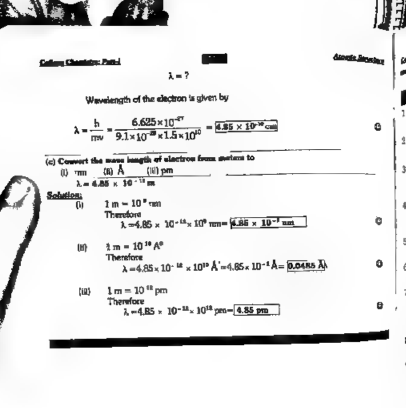
Φ

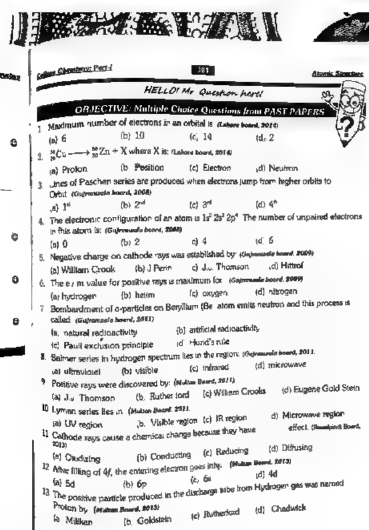
Mass of electron = m= 9 1
$$\times$$
 10 37 isg Velocity of light = c = 3 \times 10 8 m/s Velocity of electron= $v = \frac{c}{2} = \frac{3 \times 10^{6}}{2} = 1.5 \times 10^{8}$ m/s h = 6.625 \times 10 84 J s

Wavelength of the electron is given by
$$\lambda = \frac{h}{m_V} = \frac{6.625 \times 10^{-33}}{9.1 \times 10^{-31} \times 1.5 \times 10^{5}} = \frac{4.85 \times 10^{-33} \, \text{m}}{4.85 \times 10^{-33} \, \text{m}} = 4.85 \, \text{pm}$$

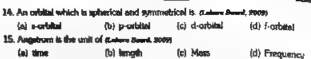
(a) Convert the mass of electron into grams and velocity of light into on e. Calculate the movelength of an electron in on.

10⁻¹² are 1 pm





Combine Park



16. Properties of waves are: (*********** a.

(a) Wave length (b) Wave number (c) Frequency (d) all

17 The nature of anode rays depend on (Ramboul Board, 2009)

(a) The nature of the electrode (b) The nature of the residual gas

(c) The nature of the discharge tube (d) All of above

18 Total number of spectral regions in a spectrum is: (Labora Board, 2010) (a) 4 (b) 6 (c) 7

19 The value of Planit's constant is (Lahou Sound, 2020)

(a) $6.62 \times 10^{-34} \text{ J.s}$ (b) $6.62 \times 10^{-27} \text{ J.s}$ (c) 6.62 × 10⁻²⁵ J.a. (d) 6.62 × 10⁻³⁵ J. s

20. In discharge take experiment the pressure of gas was measured at (Bahmalow Board mis-(a) 760 torr (b) 0.1 torr (c) 0.01 torr

21 The number of neutron present in 19K is (National Borns, 2011)

(a) 39 (ы 18 (c) 20

22 Lyman series occur in: stators Board, 2007)

(a) visible region (b) U.V region (c) IR. region (d) None of these 23 Belmer series is found in (Fatesiahed Board, 2007)

(a) L.R. region (b) U.V region (c) visible region (d) None of these 24 Which equation correctly represents the Helsenberg's uncertainty principle? (Fem. Bond, 8010) (Republic Bond, 2013)

(a) $\Delta X \times \Delta P = \frac{h}{4\pi}$ (b) $\Delta X \times \Delta P > \frac{h}{4\pi}$ (c) $\Delta X \times \Delta P \ge \frac{h}{4\pi}$ (d) $\Delta X \times \Delta P \leq -$

25 ^M_PCu + ¹_An → ^M_ACu + "x" What is "X"? (Falselebed Board, 2010)

(a) electron (b) proton (c) Bela rays (d) gamma raya The wavelength of Lyman series lies in the region. (Bergodie Board, \$010)

Ut U.V (b) visible (d) None of the above (e) 1.R

AD HEREL WAR

27. Number of neutrons present to $^{12}_{11}$ K is (D.G. Rhen Seers) 2010)

(6) 19 (c) 39

300

Walte of the Reciberg's constant is (0.0. Nam Board, 1916)

(a) 1.7904 × 10³ m⁻¹

(b) 1 9768 x 10² m⁻¹

(d _09678 × 10⁷ m⁻¹

(d) 1.6 x 10² m⁻¹

Mass of electron is (Lations Board, \$0.11)

(a) 9.1095 × 10³¹ kg

(b) 9.1096 × 10⁻¹⁶ log

(c) 9,1095 × 10⁻²⁷ kg

kl) 9 1095 × 10⁻¹¹g

M. Neutron was discovered by (Labor Found, 2011) (Surposte Board, 2014)

(a) Chadwick (b) C.D Anderson (c, Rutherford (d) Goldstein

III. When 4s orbital is complete, the electron goes into (8-3-4). Here & 2010)

tal 4p orbital (b) 3d (c) 4d

(d) 4f

The limiting line of Balmer Series lies in the region (Response Seed, 2012) (a) visible (b) U.V (c) Near I.R.

3. Lyman Series lies in spectral region (Sarpodia Board, 2012)

(a) Infrared (b) ultraviolet (c) visible (d) none of these

	Ansu	ers to l	Autiple	Charce	Questia	us Jena	Past Pe	quers.	
Q#	Ana	Q#	Ana	Q#	Ana	Q#	Ana	Q#	Ame
1	(d)	2	(c)	3	(c)	4	(b)	- 5	(c)
-	_ (a)	7	(b)	8	(b)	9	(d)	10	(a)
12	(c)	12	[p]	13	16,	14	(a)	15	(6)
1 <u>b</u>	(d)	17	(6)	18	(c)	19	(a)	20	(c)
-21	[c]	22	(b)	22	(c)	24	(c)	25	(d)
- 50	_(a)	27	,a)	28	(c)	29	fb.	30	,ů
41	_ (b)	32	(b)	33	(b)				

Detailed Explanation of Past Papers MCQs & unswers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK-I

ODE RAYS AND ITS PROPERTIES

Testisting
This will you prove that cathode mys are material perficient with negative charge? (Labora
Rend, 2007) Gulmanesia Beand, 5006; Malter Beand, 5007
Solfrede mys have reducing effect for can cause chemical change)? (8.4 New Seend,
5010 Matter and

of Contract Party

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Cook Williams

(3) How cathode rups are sermed as electrons? (Future

(1) Discuss properties of eathode roys. Find Autor Second 20121

POSITIVE RAYS AND ITS PROPERTIES

(1) Which observation tells the presence of positive rays in discharge tube? (Fairable of

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- 2000: D.G. Rhos Board. 2009)
 (2) Give reason for the production of positive rays: (Labors Board, 2013)
- (3) Write properties of positive rays (Goponicolo Board, 2011 Labore Board, 2013)
- (4) The elm value of Positive Rays is less than Cathode Rays. Justify (Madron Board, 2015)
- (5) Explain the experiment which help us to understand the discovery of protons, given

NEUTRON AND ITS PROPERTIES

Short Overtions

- What particles are formed by the decay of free neutrons? Write balanced are fractional Board, 2009; Surgedia Board, 2007, 2009, 2013. Labora Board, 2014)
 Write down mackets reactions involved in the conversion of Cu (nto Zn. (Gapanala))
- 2016) Out How the emission of a β -particle results in the increase of atomic number of element? (Randpheli Band, 2013) [3] Complete (Fatalahat Band, 2011)

- fall $\frac{1}{2}He + \frac{1}{2}Ba \rightarrow \frac{2}{2}$ (b) $\frac{1}{2}H + \frac{1}{2}H \rightarrow \frac{2}{2}$ (4) How neutrons are used in the treatment of concer? afjeur Board, 2012)
- alpindi Board, 2009: La (5) Write two properties of neutron. (Generale Board, 2009: Real . 1558
- (6) Write balanced equation for any two nuclear reactions youturn Bosta, 2008. Pd 20131

Committee

- (1) Describe the discovery and Properties of neutron in Chadwick experiment. (III.) Bourd, 2010; Charamonia Bourd, 2011. Maiten Sound, 2012)
- (2) Discuss Chaduick's experiment for the discovery of neutron. Compare the prop electron and proton. (Rundpink board 2007; Mukan board 2007)

MEASUREMENT OF CHARGE TO MASS SATIO, CHARGE AND MASS OF ELECTRON

 Colculate the mass of an electron when elm = 1.7568 × 10° C kg² tts 2014. Feliablish Board, 2011. Notices Board, 2011. 2019, OR Hote the mass of electron is Colculated by using e/m volue? (Molcon Bound, 2010; Bohmsolper Board, 2011 Labort Bot Bound, 2014)

- Law Constitute
 (1) How etn (charge to main ratio) value of electron was measured. (849)
- (2) Explain Millian's oil drop experiment to determine the charge of an election (nde Bourd, 2012: Labors Board, 2013: Labors Board, 2014. 4. 2010 Guda

STREFORD ATOMIC MODEL

the state of nucleum of the state of surface of surface of nucleum of the distri? (Supramode Hours, 2008)

What are the defects in Rutherford's atomic model? (Rund) of, 2009; No. 2009: Foliadabed Bound, 2909. 2013. Outranade Board, 2023) Give two defects in guer and a superior of the sup

Der Hors Write dejects in Rutherford's model of atom. How Bohr removed them? always and #113

UNICK'S QUANTUM THEORY, WAVELENGTH, FREQUENCY, WAVENUMBER

-) Give postulates of Planck's theory. OR What is Plank's theory mandatus sours, 2009: Labore Board, 2013)
- (i) Derive the formula for frequency of photon (only in two steps) 40.6. Elem Board, 1919. Differentials between (or What is frequency and wave number (p.s.). Khas Board, 2010.
 Multan Board, 2010.
 California Board, 2011.
- i) Gue the relationship between energy and frequency. (844)
- The energy associated with violet colour is greater than red colour in visible spectra. Why? (Lahore Sound, 2007)

as Guerians $\mathbb{R}[0]$ Wite three points of Planck's quantum theory, (ii) define frequency and wavelength. Calculated Board, 2020)

TULATES OF BOHR'S ATOMIC MODEL, RADIUS OF ORBIT

- Why the electrons move foster in an orbit of smaller radius? (Serpote & hose do you come to know that the velocities of electron in higher orbits are less than those of lower orbits.
- The radius of first orbit of hydrogen atom is 0.529Å. Calculate the radius of 3rd orbit of hydrogen atom. (Outromals Board, 2013)
- Dutify that the distance gaps between different artists go on increasing from the lower to the higher orbits. (Poteolobus Board, 2007; Sargodia Board, 2000)
- Chattings

 Chattings

 Chattings

 Delte radius of revolving electron in n-th orbit of an atom. (Gap-wassie Board, 2008, 2009, 2009).

 Delte radius of revolving electron in n-th orbit of an atom. (Gap-wassie Board, 2008, 2009).

 Sandabad Board, 2019, Behanselper Board 2009; Sahnselper Board, 2019.

 Sandabad Board, 2012, 2013).

 Gibe Deard, 2012, 2013.
- Ging Postulates of Bohr's atomic model. (Labora Baard, 2009; D.G. Rhen Board, 2011 94 Sound, 2012; Schennelpur Board, 201

BROY OF ELECTRON (BOHR'S ATORIC MODEL,

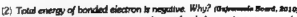
Why potential grieng, of an electron lot bonded electron) is negative in an orbit of atom?

Describe grieng, of an electron lot bonded electron) is negative in an orbit of atom?









- (3) The energy, difference between adjacent levels in an atom goes on decreasing straph Why? (Busshand Sand, 2007)
- (4) Colculate sonisation energy of hydrogen atom by using Bohr's atomic model, the

Long Countless

(1, Derive the formula for calculating the energy of an electron in 1th orbit using Bob. model: (Patabled Borri, 1971). And Karlest Borri, 2012)

SPECTAUM

- (1, Define spectrum. Name its two types. (D.G. Khan Board, 2012; Makes Board, 2007, 2009; Lab., at many on What is spectrum? Give one example. (Belausher Beert, 2009)
- (2) Why atomic spectrum is line spectrum? (Labora Boord, 2010)
- (3) What is alterric emission spectrum? (Surpodia Board, 2014) OR What is the origin of the granion of line spectrum of an atom? (Receipted Board, 2018)
- (4) What is the origin of line spectrum? (Soundle Road, 2009)
- (5) Differentiate between line spectrum and continuous spectrum. (D.G. Rion Bowd, 1018; Labor Bowd, 2014; Supplie Bowd, 2011. Nation Bowd, 2012)
 (6) Differentiate between atomic emission and atomic obscription spectrum (Behavior Bowd)

- (1) Describe atomic emission and atomic absorption spectrum with diagram. A
- (2) What is specimen? Differentiate between continuous and fine spectrum. (0.6. Number)
- (3) Define Spectrum. Explain extensic evolution and atomic absorption spectrum with djam Samel, 2023)

BOW BECYNUM OF HYDROGEN ARCM, EXPLANATION BY BOHR'S THEORY

- What is the origin of hydrogen spectrum? Ø.6. Shee Beert, 2019.
- (2) Write names of spectral series of hydrogen spectrum, assure mural, \$41.1)
- (3) What is Lymon series? In which region it lies? (Rossa, Aug. 2011)
- (4) What is the origin of hydrogen spectrum on the book of Bole's model?
- 🔯 Write down the equation for energy difference of two orbits of H-atom (Feb

DEFECTS OF BOHE'S ATCRICE MODES,

- (1) What are the defects of Bohe's atomic model 4th 9 is. Shan hand, 2009; OB Give two dejects of Bohr's atomic model. (D.G. Khan ben't, 397) Nahambanda Band, 2011. Palabanda Band, 397: Majan Band, 397: Majan Band, 397: Majan Band, 2011. Palabanda Band, 3982: Majanda Band, 2011, 2010; NOTE: For short quality, two dejects will be adopt available. White Street, 2919. Principled Point, 2912. Beginne source, 2913, 2919, 2919, see Affect will be what projecting

 (2) What is H_o line in hydrogen spectrum? Which effect explain these lines? (0.0.10)

Charlette: Patel



9810: Beleitselper Bound, 2010, 2011)

(3) Differentiate/Describe/Define Stark and Zeeman effects (Feetable Seed, 1911, Con-

- Differ artifacts of the Board, 2011 in a Zeither effects (Panished Board, 2009, 2010: D.G. Klein Board, 2011 in a Zeither Board, 2012)
 State Zeerman effect OR What is Zeitherine Board, 2012)
 D.G. Klein Board, 2006. Multim Board, 2006, 2013; Sargadha Board, 2013, 2013) Question!
- (1) Gios defects of Bohr's Atomic Model. (Labor Bons, 2011) Of Describe defects in Bohr's plomic model. (Multiur Board, 2009, \$018; Labore Board, 2010)

X.RAYS, MOSELEY'S LAW

Short Ownerforce

- 1) What are X rays? How they are produced? Supple Board, 2007: 0.6. Shee Board, 2007: Francisco Boord, 2012)
- (2) How the K-series, L-series and M-series of X-rays spectrum are produced? Assertes
- B) What is Moseley's law? Write importunce of Moseley's law # Rham Bouled, 2010: Makina Board, 2010: Labore Borrd, 2010; can Give Importance of Moseley line Planet Kenturals Bound, 2017: Resembled Bound, 2013; OB Give the Monteley's equation. Also write its importance (c" significance) (Formblad Board, 2007; Labore Board, 2008; Mahou Board, 2012)
- (4) Define Moseley's face. Give its mathematical expression. Online Board, 2001: Capurd, 2009: D.G. Klein Board, 2012: Falmitchal Berrit, 2012: Singedie Steed, 2010, 2010; Oir White is Moscley's (co.), (Falestobel Barrel, 2008; Latino Board, 2011, 2012; D.S. 47nn Board, 2009)
- (1) What are X-rays? Give the conclusions drawn by Moodey from the study of spectral fines. (Februlabed Board, 2015)

DE-BROGLIE'S EQUATION, HEIGERBERG'S UNCERTAINTY PRINCIPLE, ORBITAL

- Short Disputeres

 (1) Write and explain de Broglie's equation. States from 2010.
- A Electron has its dual nature, Justify, (5-godin Board, 2011)
- State Heisenberg's uncertainty principle and write down in mall form/equation/formula (Gapman's Band, 2007 Faintier) Band, 2015, Make Band, 2010, 2011, 2015, Behavior Band, 2007; Labor Band, 2007, 2015, 20171
- Al Equipm orbital (Labore Search 2016)

BOOK OF SEQUATION, RESERVED OF UNCERTAINTY PROCESSES OF CHARACTERS

- State Comparisons

 13 What is the function of principle quantum number? Administration of principle quantum number?

 2 Note: Surgedia Board, 2013)
- Define quantum numbers. What is the significance of Asimathol quantum number. (D.A.

 Number and also its irreorderes. The Board, 2012: On Define Asimuthal quantum Number and give its importance.

 3) State spin quantum rumber(s) briefly, dubor floor, 2013. State floor, 2011.
- (1) Describe shapes of a and p orbitals (Subsequent South State St

(1 What are quartum numbers? Give the eignificance of any one quantum number as,a, then Awal, 2009; (2) What are quantum numbers? Discuss their significance? (Principled Board 2007: Birman (3) Discuss (i) Astrochol quantum number (ii) Magnetic quantum number (superline to (4) What are quantum numbers? Discuss Principal and Azimuthal quantum numbers. (5) What are quantum numbers? Explain Principal and Magnetic quantum numbers. Other Brand, 2011) (6) Draw and explain shapes of s- and p-orbitals. (Reveloted Bowl, 2009) 7) Define orbital. Discuss shopeS of its types, (Lahore Board, 2010) ALFBAU PRINCIPLE, PAULTS EXCLUSION PRINCIPLE, HUND'S BLILE When is Author principle? (Behavelow Boors, 2008) On Define and explain n+1 ride pure Sound, 2012), ON Why 4s sub-shell is filled first and 3d afterward. (Surgodia Bound, 2013) 72, State Paulis Exclusion principle with an example (Labore Board, 2007, 2011, 2004, 2013, 2014) (3) State Aufhau principle and Pauli's exchasion principle. (Lohorz Bount, 2010) Define: Describe Hund's rule. Explain with extraple. (Anal Kadust: Board, 2012; D.G. Rhos Board, 2010; Sargedin Board, 2012; Labors Board, 2014; Fatadabad Board, 2013; Multan Board, 2018.
 Define: State Hund's rule and Paulis Exclusion principle (Convenuels Board, 2011; D.G. Board). then Resed, 2012: Resembled Resed, 2013) (1) Explain the following rules with examples: (i) Pauli's Exclusion principle: (ii) Hund's ELECTROPIC CONFIGURATIONS OF ELEMENTS Short Questions Write electronic configuration for an element with atomic number Z=29 (Families) Secret, 20111 (Z) Give the electronic configuration of Ca₂₀ and Br₂₆ (Februarios Bourd, 2009: D.O. River Bourd. ALC: U (3) Distribute the electrons in Cis₂₀ and Br₂₀ (Fatinished Board, 2008; Revealphot Board, 2011)
 (4) Write electronic configuration of 19K and 20Cu OR Write electronic configuration of

elements with atomic number 19 and 29 (Lahors Board, 2008: Feledabad Board, 2015)

(6) Write electronic configuration of elements with atomic number z= 24, 2=37 (Sorgalian Board, 2012, 2013)

(5) Write down the electronic configuration of Fe(26) and Br(35) (Labore Board, 2014)

0.08

College Chambers, Part-1

神論意。

Administration of the last of

Colors Chemistre Ports 335 TEST YOUR SKILLS Marks 97 OBJECTIVE per: 20 Minutes Der weiting, cutting, ezesing, using lead penal self result in loss of marks.

Dier weiting has four possible answers. Cheese the correct enemes.

The serve innote of armon Series has in the Hade: 17 The very sample of syman Series him in the region

(b) Varies ø (c) 1.R (d) Nove of above How many electrons can be accommodated in a sub-shell for which $n=3,\ell=1$? (b) 6 (c) 18 | C| .8 | dl .32 | The electronic configuration of an electron is 1e³, 2e⁴ 2pt | The number of supplied electrons in this alices is | (a) 2| | (b) 4| | (d) 5| | (d) 6| | (e) 4| | (e) 4| | (e) 4| | (e) 4| | (e) 6| | (e) 7| | (e) 8| | (a) 8 (d) 32 瞬 340 Positives jons are formed from the natital stom by the loss of (a) Positives: (b) protons (c) electrons (d) nation нέ (a) Positions (b) protons (c) electrons (d) neurons Cathode stys are deflected by (b) a magnetic field only (c) an electric field only (b) a magnetic field only (c) by both Ni (d) by none whe for positive rays is maximum for Eutropen (b) Heliza Nij (c) Osygen A) Häydrogen
 Neutron possess (d) Nemous |a) positive charge | (b) negative charge | (c) no charge | (d) all are connected. Lines of Peachers series are produced when electrons large from higher orbits to o |a) 1" | (b) 2" | (c) 3" | (d) 4" When atoms are votatilized, they form (a) continuous spectrum (b) line spectrum (c) electromagnetic spectrum (d) none in Milbison method for determined on of charge or electron the air th the chamber is united by (a) Protons (b) Electron Seid (c) X-rays (d) O -particles Swallest charge of electricity that has been measured to fit a (a) charge on ⁽²⁾ -porticies (b) charge on electron (c) charge on Kestya (d) charge on garriera etya particles resembles

a) He**

(b) He* (c) He more (d) He, molecular Splitting of speakers it times of the hydrogen stands under the influence of electric field is called a Salve effect (c) Compton effect (d) Photographic effect (d) S.S.S.S.A. (e) 0.5.S.S.A. (e) 0.5.S.S.A. (e) 0.5.S.S.A. (e) 0.5.S.S.A. (e) 1.5.S.A. (d) He molecule (c) He atom hig (a) Missemberg's principle (b) Auf beu principle (c) Herd's Raie (d) Pauli exclusion principle
Paucher, bracket and P und sense for a the ich Microueve region (a) visible zogazn (b) Ultraviolet region SUBJECTIVE That, 2:1D Magne Super East Section 2.3 and 4. Write any TWENTY TWO(23) short ensures. While writing answers write Wallow of Questions 2.3 and 4. Write any TWENTY TWO(23) short ensures. While writing answers write Wallow reform numbers capefully.

Sections

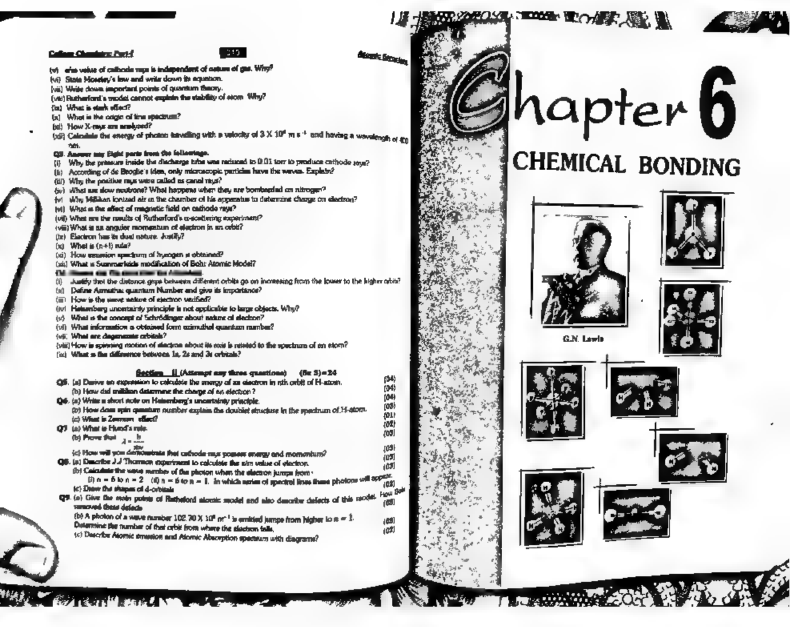
Assume any Pight parts from the followings:

Definentiate between line spectrum and continuous spectrum.

How positive rays are produced?

Define elemic orbites what is about the probability of finding electron between two orbitals?

What is the sauce of origin of x-rays?





CONTENTS

Chapter-6

e e cal Bonding

INTRODUCTION

Cause of chemical combination Energetics of bond formation ATOMIC SIZES ionic radii and covalent radii IONIZATION ENERGY, ELECTRON AFFINITY AND ELECTRONEGATIVITY

Ionization energy Electron affinity Electronegativity TYPES OF CHEMICAL BOND

Lewis concept Modern theories of covalent bonding

Valence shell electron pair repulsion theory (VSEPR)
Valence bond theory (VBT)
Atomic orbital hybridization and shapes of molecules
filocular orbital theory (MOT)

BOND ENERGY, BOND LENGTH AND

DIPOLE MOMENT Bond energy (Bond enthalpy)

ionic character and bond energy

Bond length

Dipole moment

Dipole moments and molecular structure

EFFECTS OF BONDING ON THE

PHYSICAL PROPERTIES OF COMPOUNDS

Objective and short answer, questions (exercise)

Past Papers MCQs and Short Questions

Test your skills



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INTRODUCTION

Chemical Bond

A force that holds two or more stants or loss together in a compound is called a chemical bond.

OUTET RULE: Course of Chemical Combination

Octet Rule

The tendency of atoms to attain a maximum of eight electrons in the valence shell is lmown as octat rule

Examples.

(ii) $_{12}Mg~(1s^2~2s^2~2p^6~3s^3)$ loses two electrons to form Mg^{*3} ion $(1s^3~2s^3~2p^6)$

(III) _oF 1s⁴2s²2p_p²2p_p¹2p_p²1 gain one electron to form F¹ (cm. 1s⁴2s²2p_p²2p_p¹2p_p²)

Explanation. (way arous союжь тоогичег?)

 G.N. Lewis and W. Kossel, studied the electronic configurations of elements in both free and combined state. They observed that Noble gaves are highly mert. They do not read They form only few compounds e.g. XeF₀, XeF₄, XeOF₆, XeO₅ etc. A noble gas does Dot react with another noble das.

it is because noble gases have eight electrons roctet) in their valence shell. Their general electronic configuration is 1s⁶ (He) or res² np⁶ (Ne. Ar. Kr. Xe. etc.)

Examples

_jHe 1s³

26Ne 1sh 2sh 2ph

Thus, octet of noble gases is complete. Hence these are stable. All other elements also have tendency to stabilize themselves. An element can stability by losing gaining or sharing of each one with other elements to get noble gas configuration



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terminal libraries

NOTE:

- Many compounds do not obey octet rule.
 - e.g. Formation of PCI₆ SF₆, BCI₉ etc. Hence, octet rule is not universe,
- The losing, gaining or sharing of e⁻ by an atom depends upon the conductors
 - e.g. In the formation of NaH, H accepts an effrom Na.

However, in the formation of HF. H donates major shares of its electron to the F

Lucryctics of Bond Formation.

According to modern theory of chemical bonding, aloms form bonds because if decreases the energy.

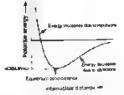
Example

Consider the formation of H_2 molecule by two hydrogen atoms.

Development of Attractive and Repulsive forces

- When two atoms come close to each other, both attractive and repulsive forces are developed between them simultaneously
- Attractive forces bring the alone close to each other and potential energy of the system is decreased
- Repulsive forces push the atoms away and potential energy of the system is increased.

The magnitude of potential energy for attractive forces is more than for repulsive forces. Thus, potential energy of the two hydrogen atoms decreases when they come close to each other.



Compromise Distance or Bond Distance

in H₂ molecule, the potential energy of the system is at minimum at a distance of 75.4 pm. At this point, attractive forces dominate the repulsive forces and two hydrogen atoms are bonded to each other. Thus, at is called the bond length or bong distance of compremise distance of two hydrogen atoms.

d Formation Energy

At compromise distance, two hydrogen atoms are at a distance of minimum energy and ese are maximum stabilized. The amount of energy released is during formation of the molecule is 436.45 kd mol it is called bond formation energy

If repulsive forces dominate attractive forces, the energy of the system received mises instability. Thus a bond is not formed

Callers Charriette: Part-I

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TOMIC SIZES

To understand bonding, relative sizes of atoms should be known because many physical and charactal properties are related to it

The sizes of atoms are expressed in terms of atomic radii, ionic radii, covalent radii etc depending upon the type of compound used for its measurement.

ATOMIC RADII

The average distance between the nucleus of an atom and the autormost electronic sell white considering it spherical is called atomic radius.

The atomic radii are usually measured in picometer (pm)

 $1 \text{ pm} = 10^{-12} \text{ m}$

Marsin carraits

- Atomic radii cannot be determined directly due to following reasons.
 - 1 There is no sharp boundary of an atom. The probability of finding an electron never becomes exactly zero even at large distance from nucleus
 - The electronic probability distribution is affected by nelabbouring atoms. Hence, size of an atom is changed from compound to compound
- Atomic radii are measured from the distance between the centres of two adjacent atoms. The distance between atoms is measured with the help of X-rays or by spectroscopy

to facilities to

Along Period

- filamic redil decrease along the period from left to right due to increase in positive charge on the nucleus. However, number of shells remains same along period.
- Due to increasing nuclear charge, nucleus powerfully attracts the resulting in the decrease of atomic radii. Shielding effect remains same from left to tight.
- In transition elements, the decrease is small from left to right due to intervening (inner) decirons e.g., from Sc(21 to Zn(30) and Y(39) in Cd(48)

Atomic radii increase down the group

- It is due to two factors
- II) Increase in Number of Shells (II) Increase in Shielding effect
- Increase in Number of Shells

In a group, number of shells increase downward. Hence, alomic radii increase









The docrama in force of attract one in culted shielding effect. e so inner alreft at

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- Econom that group, numbers of shells increase. Thus, outer electrons are less straight by micleus due to studding effect of wher shell electrons. These outer electron therefore, move outward & honce atomic radii increase.
- The decrease is more in second period and jess in higher periods.

ONIC RADII

The reduc of the too while considering it spherical to called lorde radius.



These are generally measured in <u>picometer</u> (1 pm = 10⁻¹⁰ m)

An agent may lose or gain electrons to form positive or negative ions respectively

An atom loses one or more electrons to form cations.

- fonic radius of a cation is denoted by r.
- Positive ion is please smaller in size their their parent atom.

It is due to two reasons

- (0) in positive ion, number of electrons is reduced but positive charge on nucleus renex same. Therefore nucleus powerfully attracts outer ejectrons inward resulting to decrease of sonic radius
- (M)Sometimes, outer shell is also tost due to removal of electron, therefore in decreases.
- The funic rudius of cation decreases with increase in positive charge (effects nuclear charge) on the ion.

Generally greater the positive charge, smaller the size of ion and vice yersa

It is because due to successive four of electrons, the nuclear charge attacts remain electrons more powerfully and hence size is much decreased.

Thus, the decreme in radius is larger for divident ions e.g. Mg** and still larger trivalent ions e.g. Alf-

Examples:

Ne etom = 186 pm Na *ion = 95 pm Be atom = 90 pm. Be* ion= 31 pm

R Ar of the gather bear Anica.

- An atom gains one or more electrons to form anion
- basic radius of an anion a denoted by r
- Regardon loss in advances forger in other than the pureet exam

t is due to two reasons

- # Electron-electron repulsions increases due to increase in number of electrons in the valence shell. Thus ronic size increases.
- gr) Due to increase in number of electrons, the hold of nucleus on electrons decreases Therefore, electrons move away, and thus ionic size increases.
- The lorde radius of negative into incremes with the increase in negative charge on

Exemples.

Clatom = 99 pm

Cl* ion = 181 pm

0 atom = 66 pm

Of - ion = 140 pm

to stem or Trend is Periodic Table

The variation is similar to atomic radius.

in a Group

lonic radii increases down the group due to increase in number of shells and shielding diect.

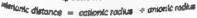
Alone Period

lonic radii decreases from left to right due to increase in nuclear charge

fonic radii for metals are for positive ions and for elements of group VA to VIIA are tox negative some.

iner-louis Dista

The Inter-tunic distance between two loss in a crystal in to the sum of outlante and anionic redus-



+ 6 - (1) F.

Pauling determined 'PI' for KCI crystal as 314 pm. II is lettedly the sum of radu of K* and CI ions









Company Profes

718



R = 133 + 181 = 314 pm

it shows that ionic radius is an additive property.

Re-arranging equation (1) , τ of various ions in potassium saits can be determined as

Similarly r_{τ} can also be determined in various chloride saits as

$$r_{\star}=R\cdot r_{th}.$$

COCAH STROOM

The expelent radius of an element is half of the single bond length between to populatily bonded in a molecule.

Expression:

). The band length of $H_{\rm g}$ molecule is 75.4 pm. So, cavalent radius of H# 75.4/2 = 37 7 pm

 The bond length of Cl₂ molecule is 198.8 pm. So, covalent radius of Cl is 198.8/2 = 99.4 pm.



The consist rodius of one atom can be used to measure the consient radius of quother atom.

. . H .d. 47

Bond length of C - Cl bond in CH₈Cl is 176.7 pm. Since covalent radius of Cl is 99.9 pm, famelore covalent radius of C should be 176.7 -99.4 = 77.3 pm.

The partialism of covalent radii in periodic table is almost similar to atomic radii

trasportion Emergies, Libertron Alfinity and Electronequitivity

Since band formation involves energy changes, therefore, thermodynamic property must be known to undentand chemical bonding.

These properties are ionization energy, electron affinity, electronegativity ex-

NZAHON ENERGY OR IONIZATION POTENTIAL

the ionization energy of an element is the minimum energy rec the from its gaseous atom to form as los,

g is sometimes called as lontzation Potential.

it is expressed led/mot or electron volts (eV)/atom

1 eV/etom = 96.48 kJ/mol

 $Mg_{igi} \longrightarrow Mg_{igi}^* + e^*$ $\Delta H = + 738 \text{ kJ/mole}$

was olering ionization energies

Atomic radius of the atom

increase in atomic radius decreases the ionization surryy. It is because when size sureases, the hold of nucleus on outer electron discreases. So, less energy is required to remove them. Hence, ionization energy decreases.

Shielding effect of inner electrons

increase in abornic radius decreases the ionization energy. It is because when shielding effect increases, the hold of nucleus on outer electron decreases. So, less energy is required to remove them. Hence, sonization energy decreases

libelear charge or proton number of the atom

Increase in nuclear charge increases the sonization energy it is because when nuclear charge increases, the hold of nucleus on outer electron also increases. So, more energy is required to remove them. Hence, unication energy increases.

Nature of orbital

The s-orbital is small and compact so it is difficult to restore electron from this orbital.

The Forbital is sarger in size, therefore, it is easy to response electron from this gebital.

Hence, order of decreasing ionization energy is a>p>d>1

house front in Periodic Tolds

of Glond

totalen energy decreases down the group attrough nuclear charge increases

he due to increase in number of shells and shielding effect down the group

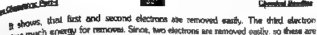
Due to increase in number of shells and shelding exect quows are property to addition of shells, electrons become away from nucleus and lorde of attraction becomes away from nucleus and lorde of attraction. Sheeding between rescieus and outer electron decreases due to increase in distance. Sheetding





longation energy decreases.

effect of issuer electrons also increases. Hence outer electron is easy to remove



is structured to restroyed. Since, two electrons are removed easily, so these are de valence electrons.

Alone Period

- longation energy increases along the period from left to right.
- It is due to increase in nuclear charge along the period. Along period, proton nur excesses and thus positive charge on the nucleus increases. This high nuclear du powerfully attracts electron. Thus, more energy is required to remove the electron therefore, contraction energies increases.
- Each period starts with an element, which has one electron in its valence shell and with the completion of volume shell.

Michael Imposition Energies

- When first electron is removed from an atom, ionization energy is known as i CONTRACTOR CONTRACTOR
- When second or third electrons are removed, these are called Second ionization or or Third contration energy respectively and so on.
- Plast, second & third contration energy values increases in the following order
 - First I.E. < Second I.E. < Third I.E.

It is because, due to removel of electron, hold of the nucleus on the remaining ele increases. Hence, they are difficult to remove and thus ionization energy increas

Emmoles → Mg_m*+ e* Mg_{et} 1 Mg_M → Mg_M + e

4 H =+ 738 ki/mole

Mg_{al}²⁺
$$\longrightarrow$$
 Mg_{al}³⁺+ e'

 $\Delta H = + 1451 \, kJ/mole$ ∆ H = + 7730 kJ/mole

Suplications Of Jones Con Factor

- Indication of metallic character Generally
 - Elements with low contration energies are metals.
 - Elements with <u>high ionization energies</u> are <u>non-metals</u>
 - Elements with <u>intermediate</u> values of ionization energies are <u>metalloids</u>

2 Indication of valence electrons

lonization energy measurements provide evidence for the number of valence significant Example

For magnesium, the ionization energies are

First I.E. = +738 k.l/mot Second I.E. = +1451 k.l/mol, Third I.E. = +7730 k.l/m²

- inflortion of Noture of Compounds Demonts with low rontzation energy values (e.g. group I A and II A), can easily give electrons, hence, they form jonic compounds.
 - Elements with intermediate ionization energies form covalent compounds by sharing electrons with other elements.

Sobility of Atom

- . In gaseous phase, atoms and ions are isolated and have no external influences. So, tonization energy gives a quantitative measure of the stability of squated atom.
- Generally, higher the lonization energy, higher the stability of aroms, e.g., Noble gases have highest contration energies in their respective periods, so they are more

DECTRON AFFINITY

The electron affinity of an atom is the amount of energy released when an electron adds is an ainsty or partially filled orbital of an atom in the releases shall to form an autom ring a unit negative charge.

It is expressed as kil/male.

Emmple:

Electron affinity is the measure of the attraction of the nucleus of an atom for extra electron. Clus

lober Electron Alfantics

- During addition of first electron, energy is released, therefore, electron affinity is given Tegative sign.
- --> O'₀₆ ΔH ≈ -141 κΔ(mo) When a second electron is added in a uni-negative ion, the incoming electron is is realised to the by the negative ion and energy is absorbed in the process.
- O₁₀ + e → O³ to AH = +844 kJ/mox Phus that electron affinity of an atom is negative, while second, third etc electron affinities are positive.



College Chambring Part I

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Direct measurement of Electron affinity is difficult. Therefore. Electron affinities generally measured indirectly

Factors Affecting Lies from Affants

- Electron affinity depends upon atomic radius, nuclear charge, shielding effect etc.
- increase in atomic radius and shielding effect decreases the attraction between nucle and incoming electron, therefore electron affinity decreases
- Increase in nuclear charge increases the attraction between nucleus and income electron, so electron affinity increases.

term the same of

Along Period

- Generally, gjectron affinity values increase from left to right in a period due to detres in atomic radius and increase in nuclear charge.
- Thus alkali metals on the left have lowest electron affinities and halogens on the ig have <u>highest</u> electron affinities.

[п а Стоир

Electron affinity values decrease down the croup due to increase in number of shell at shielding effect.

It is because, atomic radius and shielding effect increases down the group. This decreases the force of attraction between nucleus and the incoming electrons. Here electron affinity decreases.

NOTE: Exceptions to the General Trend

- Electron affinity of F is less than C. It is because, F has very small size and seven electrons in 2s and 2p sub-shells. These electrons form a thick electronic cloud, which repels the incoming electron.
- The elements of group life. VA and VIIIA show abnormally low values in every pered. it is due to two reasons
 - 49 Groups IIA and VIIIA have completely filled orbitals, therefore, incoming electrons will have to go to higher energy orbitals, resulting in the decrease of section affinity values.
 - (iii) Group VA has half filled orbital. The repulsions between the incoming election and the electrons present in the half filled orbital results in the decrease of electron official vehicle

colors Charles to V. Ports

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CIRONEGATIVITY The tandency of an atom to attract a shared electron pair twowide treef in a molecule is called electronograticity.

Electronegativity has no unit.

14: tron: gaticity Scale

- Direct measurement of electronegativity values is not possible
- Pauling suggested an arbitrary scale from 0 to 4, as a measure of electronegativity of
- 11-16 based upon the difference between expected bond energies of normal covalent bonds of elements and experimental bond energies.
- in this scale, electronogativity of F is arbitrarily given as 4. The electronogativities of other elettrents are determined relative to F

dedictions of Pedyptic Table

Along Period

Electronegativity values increases from left to right in a periodic table, due to decrease in alomic size and increase in nuclear charge.

Electronecrativity, values decreases down the group, due to increasing reuniter of shells and shielding effect.

Esperiodic table.

- the most electronegative element is Flourine (F)
- the seast electronegative element is Francium (Fr).

the conegurate and Nature of Book

Difference of electromegativity values is an order to the polar nature of bond in a molecule

- If electronegativity difference is zero, the bond is non-polar. Thus, all the bonds. between similar atoms ere non-polar 4.4. Ha
- · All the bonds between different elements are generally polar 4.9. HCl
- il electronegativity difference of two bonded asoms is 1.7 or greater than 1.7 bond is
- If electronogativity difference of two bonded atoms is less than 1.7 bond is covalent.



- Chamleton Part-I

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Counting Barrie

(STES OF CHI MICAL BOND

There are generally three basic types of bonds.

- Ionic or Electrovalent bond.
- 2. Covalent bond
- 3 Co-ordinate covarent bond.

LEWIS CONCLET

TONIC OR ELECTROVALENT BOND

The bond formed by the complete transfer of one or more electrons from an atom with loss lossisation energy to consider exam with high electron affinity is called loss bond.

Explanation:

- In energy terms, the electropositive elements are at higher energy state while the electronegative elements are at lower energy state. This difference of energy results in the transfer of electrons from higher energy state to lower energy state.
- The atom, which loses electron form cation, while the atom, which gains electron form
- The compounds formed by the cations and anions are called ionic or electrovatent compounds.
- Generally, elements with low ionization energies (groups i-A or II-A,, react with elements of high electronogativities and high electron affinities (e.g. Group VI-A or VII-A elements) to form tonic bond, e.g. NaCl, KCl etc.

Example: KCl

Formation of K* ton

Electronic configuration of K is

- It can be represented as K (2, 8, 8, 1)
- K tends to loose its outermost electron and form K* Ion. The energy required to remove one electron from K is equal to its first ionization energy.

$$K(2, 8, 8, 1) \longrightarrow K^*(2, 8, 8) + 1e^- \Delta H = 439 \text{ kJ/mol}$$
[Arl

After loosing electron, potastism attain the nearest noble gas configuration of Art2, 8 8

Formation of CP1 ion

Electronic configuration of Cliatom is

1st 2st 2pt 3st 3pt

If can be represented as CI (2,8,7)

Charles, Part-I

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It tends to gain one electron to attain the pearest noble was configuration of Ar [2, 8, 8] and 349 kJ/mol energy is released. This is the discron affinity of Cl.

formation of KCI

- Thus, electron is transferred from K to Cl atom to form K* and Cl ions. These ions develop electrostatic force of attraction between them and thus ronic bonds are formed.
- R and Cl ions arrange themselves to form crystal lattice and 690 kel/mot energy is relicased. The energy released during the formation of crystal lattice is called lattice

$$K^+ + CI$$
 \longrightarrow KCI $\Delta H = 690 kJ/mol$

NOTE:

- Elements of group IA (L., Na, K, Rb. Cs) are good losers of electrons
- Elements of group VIIA(F, Cl, Br. I) are good gainers of electron.
- Thus both these combine with each other and form lonic bonds.
- Similar type of bonds are also present between elements of group IIA and group VIA elements.

Emmpres:

So, the formulae are

- Alternation oxide will be ${\rm AlgO_3}$ (i.e., ${\rm AlgO_3}$ contains one in the ratio 2.3)
- Calcium exide will be CaO (i.e. CaO contains ions in the ratio 1 1;)
- Similarly, CaS and Al₂S₈ are also conic compounds to some extent.

basic nequirity and Nature of Coreposite

- Difference of electronegativity between bonded atoms can decide the nature of
- Generally 11 electronegativity difference is 1.7 or more, the bond is ionic. Thus, NoCl has 72% ionic character. Cult has 92% ionic character. Calculate 11998, ionic character.
- Calculations show that there is no bond with 100% ionis character.



Colon Chapters Paris

1.2%

Chemical Room





and Beach

2. COVALEN) BOND (FILCTRON PAIR BOND)

The bond formed by mutual sharing of electrons between two atoms is called

During covalent bond formation, an atom completes its valence shall by staring and attains nearest ineat gas configuration.

A covalent hand may be palar or non-palar

Cartesian Constead Bonds

The copolent bond, in which bonding electron pairs are equally shared by too stone is called non-polar condent bands

Exemple

Covolent bonds between like atoms is non-polar

- In such compounds distribution of electron density is uniform. Due to even distribution of charge, the bonded atoms are electrically neutral. These compounds are called now
- CCL also have equal electron density distribution. Hence it is also a non-polar compound. It is due to cancellation of all the dipoles of this molecule due to its symmetry. In CCI₄, all C-Ci bonds are polar but molecule is overall non-polar

Other Examples:

CH4. SIH4 SICL etc.

The coodent hoad in which bonding electrons are not shared equally by close is Polar Corolen Beach celled polar constant bond

- Covalent bond between unlike atoms is polar
- in such cases the shared electrons are more attracted by the element having large electronegativity it makes one end of the molecule partially, positive and other and partially positive and other and other and other and other partially positive and other and other partially positive and other and other partially positive and other and o partially, negative Such compounds are called polar compounds. In these compounds distribution of electron density is not uniform

Exemples

Hudrogen Flouride

Charlete Part

Water

H:0: # 0

Methyl Chioride

Methanol

A cocalent bond can be classified as single, double or triple.

ger kjørg provin øsen re

If covalent bond is formed by sharing of one electron pair (two electrons), it is called a single copolent bond.

: Ci- + -Ci: --- : Ci: Ci: or Ci- Ci Example

Later test of

If considert band is formed by sharing of two electron pairs (four electrons), it is called a double constent band.

<u> स्थापनंद</u>

🕅 Охудел

即 Carbon Directric

It is formed from heterostoms. But due to linear structure polar bonds cancel the effect of each other Hence, it is a non-polar compound.



Chambelon: Part-I

called a triple consient band.

Examples

Some non-metallic atoms especially C and St, mutually share their electrons with each other and thus form long chain molecules called magromolecules.

e.g. Diamond, Graphite and SiC etc.

Carbon can form wingle, double or triple bonds in alkanes, eliames and alkanes.

The compounds of C and H containing only single bonds are called alkanes

Example: Ethane

Silicon also gives at

Example Distance

Alkenes

The compounds of C and H containing double bonds are called alkenes Example: Ethene

The compounds of C and H containing triple bonds are called alkanes Example: Pthone

Coffee Chambers: Press!



COLORDINATE COVALENT BOND

eralmeter coculent band to formed between two stome when the shared pair of

- The atom which donates the electron pair is called the ' Donor '
- The atom which accepts them for bond formation is called the "Acceptor"

T. March 1.

. This bond is generally represented by an arrow pointing from donor to acceptor atom.

Exemples

jerostron of Annoonin With Br.

- A co-ordinate covalent bond is formed by the donation of electron pair form nitrogen of NH_a to B of BF_a
- NH₁ has three covalent bonds and one electron pair on nitrogen atom.
- In BF₃ , octet of B is not complete, therefore it is deficient in electrons.
- Thus N can donate the pair of electrons to the acceptor BFs and a co-ordinate covalent

The complex so formed is electrically neutral. Changes are mentioned on N and B atoms.

In some cases after bond formation, no distinction is present between a co-ordinate coolens bond and a coodent bond.

Exemples

Connation of H₂O' Inv

When an acid is discoved in water, a bond is formed between H_eO and H* ion. In this shared pair of electron is donated by O of H₂O.

All the three bonds of H_aO^+ are equal. Every bond is 33% coordinate covalent and 66% indices



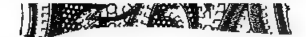
A Printed processor of the

Alcohols and ethers donate their lone pairs to H* lone to form coordinate covalent bonds. The ions produced are called exemium ion.

40 Committee of ML, Lor

A co-ordinate covariant bond is formed by the donation of electron to H* ion by nitrogen

 Like ammonia, all the primary, secondary and tertiary aminus form coordinate covatent band with H*





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PH_a combines with H⁺ to give PH_a⁺ ion called phosphonium ion.

Coordinate covalent bund are also present in HNO₃. Many oxysoids of halogens. like (HClC₁, HClO₂ HClO₄, have coordinate covalent bond between Cl and O.

I'm tations of Lewis model

model has explained how the aloms are bonded to one another and how the electron pairs are shared between the bonded atoms. Lewis model has following limitations

(f) It is an over simplified model. (ii) It does not tell about the shapes of molecules. Many physical and chamical properties of molecules depand upon three dimensional arrangement of their atoms.

(iii) It does not tell about the bond distances, various energy transitions as shown by

spectroscopic techniques.

(iv) If does not account for the unique behaviour of molecules during chemical reactions.

Goldies of an Ideal Model

An ideal model should explain molecular shapes, geometries, bond polarities, bond distincts, various enemy transitions and unique behaviour of molecules during chemical reactions

MODERN THEORIES OF COVALENT BONDING

Various theories have been proposed to explain the nature of covalent bonding.

A few important of these theories are

A. Valence shell electron pair regulation theory (VSEPR)

B. Valence bond theory (VBT,

C. Molecular orbital theory (MOT)

THE REPORT OF THE PROPERTY OF

This theory was proposed by Sidwick and Powell in 1940, to explain the shape of covalent molecules on the bases of electron pairs in the outer orbit of the central atom.

Recently, Nytholin and Sillespie developed VSEPR theory to explain the shapes of molecules for non-transition elements





pairs and bond pairs) ore arranged area non apart to keep repulations at a minimum.

% stafan vol VSEPR theory

- 1 Both lone pair and bond pairs are involved in determining the geometry of molecules
- 2. The electron pairs are arranged at a maximum distance around the central polyvalent atom to avoid repulsions.
- A lone pair occupies more space than a bond pair
- The repulsion byte electron pairs decreases in the following order Lone pair-lone pair > Lone pair-bond pair > bond pair-bond pair

It is because a bond pair is attracted by both nuclei while a ione pair is attracted by only one nucleus, therefore, lone pair is less attracted by nucleus. Hence its spread oil more in space than bond pairs and occupy more space. Thus, it exerts greater repulsions and compresses the bond pairs.

- Double bonds and triple bonds occupy more space than a single bond.
 - It is because, double and triple bonds contain higher electronic charge density therefore, they occupy more space than one electron pair of single bond.
- Multiple bands (i.e. double or triple bond) behave as single bond in determining the geometry of molecule

It is because they also occupy the same region between the nuclei like a single book

SEPTICATIONS OF VALENCE SHELL ELECTRON PAIR REPULSION THEORY

- VSPPR therap can predict the shares of covalent moterales.
- Consider a central atom A It combines with two or more B type atoms to give AB, AB, AB, AB, AB, type molecules.
- The number of B' atoms attached to M' depends upon the satency of A
- Thus, molecules can be dayled into different categories as given in the following table



	Tuble: S	ariter e	I melecul	les according to	VSEPR theory	Chambrol 6
tal No. of Electron its around atral mon	post paire	No a Tione pairs		Printingement Printingement Licetron Podra	Molecular Secondary	
2	2	1 0	AB ₂	Linear	Linear	BeXg.wdX gCug.HgCl
,	3	Û	AB,	Triongular Planor	Triengular Planar	BX,
•	2	ι	7.0,		Bent or Angeler	ShQ _p , SO ₃
	4	0			Tetrahedral	SH, CO.
	7	_			1	
4	3	1	AB _e	Tetrahedrad	Triangular Pyremidel	NH, PH, AcCl,
4	-	1 2	AB _e	Tetrahedrad		

Solecules With Two Cherman Pairs (AB, Tope -

Serrole BeCl.

There are two elactron pairs around the central Be atom.

Roth electron pairs are bond pairs

There is no lone pair According to VSLTT ..., the two electron pairs are present at angle in order to have mi . . in repursion to have This

अवतर्ग व स्टब्स् तलकात्रात्र के क्षित्रकार

The roolecular geometry is determined by bond pairs only. So Belly Rindom rodecuje is unear

The Exercise

Hard, Carly Sycle Carly Hard, Co. ex.













College Chambeles: Park-I

2 Pioles nies with three electron pairs

AB, Type: With No Lone Pate:

- EXAMPLE BE. There are three electron pairs around the central B atom.
- All three electron pairs are bond pairs.
- There is no lone pair
- According to VSEPR theory the three electron pairs are present at 120° angle in order to have minimum repulsion between them. Thus arrangement of electron pairs is triangular planer.
- The molecular geometry is determined by bond pairs only. So. BF₂ molecule is also thangular planar

Other Exemples

Hydrides of group fil-A (AIH_a , GaH_a , InH_a) and their halides(BF_a , AICl_a) etc.

II) AB. Type: WITH TWO BOND PAIRS AND ONE LONE PAIR: EXAMPLE: SHCT:

- There are three electron pairs around the central Sn atom.
- Two electron pairs are bond pairs and one electron pair is ione pair
- According to VSEPR theory, the times electron pairs are present in a triangular planer form in order to have minimum repulsion between them. Thus errongement of electrons is triangular planar.
- The molecular geometry is determined by bond pairs only. So. SnCl_a motercule is bent or angular. Since lone pair exert greater repulsions on bond pairs, therefore, $\underline{Cl} = \underline{So} = \underline{Cl}$ bond angle is less than 120°

AB. WITH MULTIPLE BONDS

in SO_a, one corner of triangle is occupied by lone pair and other two are occupied by S=O bonds. Thus it has angular structure

In SOs, all comes of triangle are occupied by S=O bonds. Thus it has triagular structure (tiangular planar).



8. States alex With Four I bernau Pales Around Central Aroun

0 AH, TYPE: WITH MO LONE PAIRS EXAMPLE: CH.

- There are four electron pains around the central C atom.
- All lour electron pairs are bond pairs.



There is no lone pair

neording to VSEPR theory the four electron pairs are present at 1035 angle in order to have minimum repulsion between them. Thus amagement of electron pairs is tetrahedra.

the molecular geometry is determined by bond pairs only. So, CH. molecule is also tetrahedral

the hydogens are present at the corner of tetrahedron. Thus CH, molecule is tetrahedral Rediti - C - H bond angle is 109.5°

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this shockure is non - planar. It has four comers, four faces, six schoes and six bond anales.

(tiller exemples:

CCL, StH, GeH, etc.

III ABLL Type: WITH THERE BOND PAIRS AND ONE LOKE PAIR EXAMPLE: NH.

There are four electron pairs around the central N atom.

Titree electron pairs are bond pairs and one electron pair is lone

According to VSEPR theory, the three electron pairs are present in a laterhedral form in order to have minimum republish between them.

This arrangement of electrons is tetrahedral.
The molecular geometry is determined by bond pairs only. So, NH

molecule is triangular pyramidal

The electronic configuration of N is

₂N 1st 2st 2p, 12p, 12p, 12p, 1

Singuione pair occupies more space than bond pain. Therefore ione pair present in 2s while of N exerts greater repulsions on bond pairs. Thus H - N - H bond andle reduces Ion 109.5° to 107.5°

Other Exemples

NFa, PCI, PHa, AsH, ShH, BiH, etc

of Estetronegatistic on Bond Ausie.

International to which hydrogens of NH₂ are subdiffused by more electronscaling stoms

a bond anales are jurther reduced.

Reason, In NFs, the paler N-F band pulls the lone pair of N closer to the F to the second of the polar N-F bond pulls the lone pain, hence bond angle is to the second of the se word to 1029 Further bonding electrons are present closer to F, due to

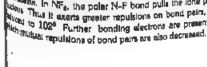






















AN ANALONS FOR THE SOME PART AND THE LOSS FROM EMPL NO

- These are four electron pears around the vertex Q alors.
- Two electron pairs are bood pairs and two electron pairs are lone pair
- According to VSSPR theory the two electron pass are present to a triangular plan down at creater it fame minimum resolution between them. Thus extended out of every A MARKETINE
- "Le raquestion geometry is determined by band pairs only So. HyO, molecule is asy **QCOBU**É
- The minimum configuration of Our

D 14 24 29, 20, 201

- fit waster than some paties and have bound prove one present. Thus, it shaes when system of septembers a losse past losse past figure gent formed past HAVE HAVE'S GAME . HAVE SHOW
- All the marker pains from total colors Total of the somework mentioners are recepted by wine power Trips that exchanges have anyther AN ISSUE SHOWS HER
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The on electron piers arrange in an actahadral

Example, 9F,

Look, whose of 25s Filt chooses

- ii). If draw not pive responsition the hormation of bond
- Hil if ribsses med eingelieben bittettelt meterophen





is based on the quantum-mechanical model of atom, it explains formation of board and entergies, board sength and shapes of mulecules

the basic assumption of V.S. theory is that

Half-filled valuence attacks orbitals of combining stress mostles to form on

The line hebreren two outles is called bond and or lines outless too

Buckleyed

- ill. The two coverlapping orbitals must be refered substate and must be not little.
- (ii) Larger the overlap etronger the bond
- hip. The America, of based is denomined by discretion of overlapping orbitals
- his Ties electrone, greenst in bood nibital most have opposite opins. Thus parced giernouse etablice (camulious

Valence hirror theory, describes two types of horid

is littled fortund i

It is the stripts bend formed, when two pertially littled elemin printeds everlap in a trip that the probability of finding the electron is maximum ensured the line justing

All single perceivent beniels gre a bund

1 termedica ad \$1, malecular

A- & Overies

The electronic configuration of H is 1st 1s orbitals of him hydrogen storin overlap with but other to form H. H bend in this band electron density is present between the nuclei This bond is called or -bond







Chronica of HIC protestale.

La Cherien

H=F bond in hormand by pairing of electrons, one from H atom and one from F atoms





Calmy Chester, Part I

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Charles .

Electronic portfigurations of H and F are

H 35°

5 16 24 2p, 2p, 2p,

Thus, H and F what our electron each to complete their outconcer shell. This is $4\alpha_{2}$ consistency of its orbital of H with $2\alpha_{1}$ orbital of F



100

Farmation of F₂ methods

4 3 Sourion

Security configuration of Fig.

F 15 25 25 25 25

Thus, $2p_a$ orbitals of true F stress constap with each other to form F-F bond.



Pi - Book of mand i

The hand formed by side-to-side or parallel courley of the p-orbitals. The hand in which greatest electron density to prevent above and below the historicalism sets.

For $\pi\text{-bond}$ formation, the two overlapping orbitals must lie in the same plane

Cornicier the x-band formation by p-orbitals.

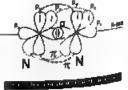
Examples, Formation of Namolecule

 N_{α} molecule is formed by combination of two strongers atoms.

The electronic configuration of N is

N 1st 2st 2p, 2p, 2p, 2p,

Two N atoms have helf-filled 2p. 2p, & 2p, orbitals 2p, orbitals of two N atoms overlap and-to-said to give a e - bond, while two 2p, orbitals and two 2p, orbitals of two N atoms overlap to give two n-bond by parallel overlapping.



TOP |

Thomas of the con-

Thus there is one σ - bond and two z - bonds between two nitrogen atoms in N_g

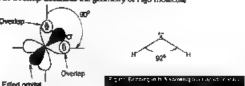
Signal bond is formed by linear head-on-overlap, while pi bond is formed by parallel godge. Therefore, in signal bond there is larger overlap than pi bond. According to Valence Bood Theory, larger the overlap stronger the bond. Thus, signal bond is stronger than pi god.

are most H 9 Meleral s

gis a none linear molecule. In this H S = H bond angle is about 92⁴. Bechools configuration of S is

25 15 25 2p, 2p, 2p, 35 3p, 3p, 3p, 3p, 3

Two iself-filled 3p orbitals of 5 overlap with two 1s orbitals of H atoms to form H₂S solecule. Thus, idea of overlap describes the geometry of H₂S molecule



Somie Orbital Hebridization and Shapes of Molecules

The process in which atomic orbitals of different energy and shape are mixed together to form a new set of equivalent orbitals of some energy and same shape is called biridization.

The orbitals formed are called behald orbitals.

Department.

Concept of intenditation is used to explain equivalent telegraderics of C, bond angles in the and NFI. atc.

for Excitation

In some cases, one electron is promoted from lower orbital to higher orbital to increase the number of unpeired electrons. The excited orbitals then undergo hybridization. The excited orbitals then undergo hybridization. The excited orbitals then undergo hybridization. The energy required for security of electron and hybridization is a simultaneous process. The energy required for security of electron and hybridization and bond immation. Hybridization gives obtain in compensated during hybridization and bond immation.



- Chartery Park!

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Types of Hybridizati

There are various types of hybridization depending upon the number and nature of orbitals taking part in hybridization e.g. sp³ sp³ sp dsp³, dsp³, d²sp³ etc

s and p orbitals hybridize in the following ways:

Hybridization.

In this, one a and three p atomic orbitals intermits to form four equivalent with hed sp3 hybrid stamic orbitals.

Four sy² hybrid orbitals forms a tetrahedral structure with mutual bond angles of 109 5



3 vor api hybrid ordale.

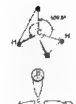
Englare up trylandization:

Example 1, CH. (Mathone)

in Clf., C undergoes ap hybridization. hybridization can be represented as

6 C (ground state) 26, 30, 20, C rescried state 24 20, 20,

C Suphridated History | A | 1 | 1 | 1 |

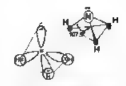


- Each ma hybrid orbital has two lobes, one smaller and one larger. Ect employed smaller tobe is not usually shown.
- Four sp⁴ hybrid orbitals of C are present at the corners of regular tetramedron.

 Four sp⁴ hybrid orbitals of C overlap with half-falled 1s orbitals of four H atoms to left 1, 109.5° Or CH, molecule with legisledia geometry. All the bond angles are of 109.50 metershedral structure of CH, has four force. tetrahedral structure of CH₄ has four faces, four corners, six edges and six bond angles.

grande 2: NHs

in Ni²s, N undergoes ap⁹ hybridization. One s and three p-orbitals of N hybridize to give ion sp hybrid orbitals. It can be supresented as



- Four sp³ hybrid orbitals are present at the corners of regular tetrahedron.
- Trees half-lifted sp[®] hybrid orbitals of N overlap with half-filled is cubitals of three H atoms to form NH₈ molecule. While a lone pair is present at one corner of a terrahedron. Thus NH₈ have pyramidal geometry in which hydrogen atoms from the base while lone nair form the apex.
- Since lone pair occupies more space than bond pairs and closer to N, therefore, essent greater repulsions on bond pairs. Thus $H \sim N H$ bond angle is reduced to 107.5° in NH $_{\rm S}$ from normal tetrehedral angle of 109.5° .

čnansie 3: H₂O

įQ.

In H₂O , O undergoes sp⁸ hybridization. One s and three p-orbitals of O hybridize to give four spr hybrid orbitals. It can be represented as







- Four up hybrid orbitals are present at the comers
- Two half-filled sp³-hybrid orbitals of O overlap with half-filled 1s orbitals of two H atoms to form H₂O molecule while two lone palls are present at the corners of a temphedron.
- Since ione pair occupies more space than bond pairs and closer to O, therefore, exerts Since ione pair occupies more space than bond pairs and closer to O, therefore, exerts since ione pair occupies more space than bond pairs. Thus H=0 H bond angle is reduced to 104.5° in the HeO from normal tetrahedral angle of 109.5°



om sp^a hybrid orbitals farms a triangular planer structure with mutual bond angles of



de L. DE,

in BF₃, B undergoes sp³ bybridization. This hybridization can be represented as

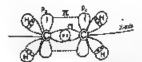
20 20, 20, 20,

-

Fluorine has one half filled p-orbital. Three sp¹-hybrid orbitals of B overlap with half-filled $2p_s$ orbitals of filters F atoms to form BF_s molecule with triangular planar structure

Example 2: Call

In C₂H₄ each C atom shows sp² hybridization. This hybridization can be represented as



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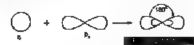
- Bach C atom undergoes sp¹-sp² overlap with each other to fown a sp bond.

 Similarly, each C also sp² is overlap with H atoms to form C₆H₆ molecule with triangular. planar structure.
- One unhybridized p-orbital of two earbon atoms overlap sideways to form a x-bond. In this bond probability of finding electron is maximum between the nuclei.

$m \cdot Hein Blocation$

In this, one s and one is atomic orbitals inhumic to form two equivalent orbitals called to induid atomic orbitals.

The two hybrid orbitals are present at an angle of 180°



Emmple 1, BeCl.

In BeCla, Be undergoes up hybridization. This hybridization can be represented as

-

0

20, 20, 29, 4 [1]

eb ab Two sp-inybrid orbitals of Be are present at an angle of 180°. These orbitals overlap with half-filled 3p, orbitals of two CI atoras to form linear BeCla molecule.

Exemple 2: Calle

In C_2H_2 each C atom shows so hybridization. This hybridization can be represented as

(i) \$0





College Chambers: Parish

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Chimical Benda

- Each C atom undergoes sp-sp overlap with each other to form a o-bond. Similarly, each C also sp-s overlaps with H atoms to form C₂H₂ molecule to form C₂H₂ molecule with linear structure
- Two unhybridized p orbitals of two carbon atoms overlap sideways to form two π bonds. Four electronic clouds of two π - bonds intermit and surround the sigma bond in the shape of a drum.

MOLECULAR ORBITAL THEORY

h is based on quantum mechanical model of atom. It considers molecule as a single unit. It's main points are

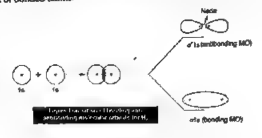
- 1 Atomic orbitals (AOs) combine to give new set of molecular orbitals (MOs), which are characteristic of the whole molecule
- The molecular orbitals surround two-or more nuclei of bonded atoms
- 3. Two atomic orbitals of combining atoms overlap with each other to give two molecular orbitals. One with lower energy is called bonding molecular orbital and the other with higher energy is called antibonding mulecular orbital
- In bonding MOs, electron density is present between two nuclei while in antibording MOs it is away from nuclei.
- The mamber of MOs produced is always equal to the number of AOs combined.
- The filling of molecular orbitals takes place according to Authau, principal, Paul's exclusion principle and Hund's rule.

Explanations - s Orealigh

Consider two hydrogen atoms H_A and H_B. Each hydrogen atom has one electron in its Is cribital. Two Is orbitals of combining hydrogen atoms overlap with each other to give two molecular orbitals of different energy.

Low energy molecular orbital is called bonding sigma MO denoted by σ is and high energy molecular orbital is called antibonding sigma MO denoted byo 1s.

The bonding molecular orbital is symmetrical about the axis (molecular axis) joining the nuclei of bonded atoms.



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e - parech p

Consider the overlap of p orbitals. Three p - orbitals are perpendicular to each other. Following overlap occurs between positizals,

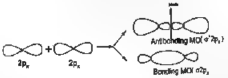
... Hand ! Head On Approach).

The MO, which is symmetrical shout the band acts is called a - MO

The MO produced by the head on everlap of atomic orbitals are called a - MO

When atomic orbitals combine head on o bonding MO and s "- antibonding MO are produced

Head on overlap of two $2p_s$ orbitals produces σ and σ MOs.



Tapet Casterna condition of the

:- 510 (Siderens Approach).

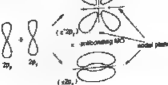
The MO, which is formed by sidewors overlap of atomic orbitals and contains only one nadal plane on the nuclear axis is colled a - MO

Ronding molecular molecular orbital $\pi\left(2p_{\mu}\right)$ and $\pi\left(2p_{\mu}\right)$ have zero electron density on the nuclear axis. It is called nodal plane. Electron density is uniformly distributed above and anelo iston ant wolled

Anthonding molecular orbitals $\pi^*(2p_a)$ and $\pi^*(2p_a)$ have least electron density in the π

Since $2p_{\pi}$ and $2p_{\pi}$ are degenerate, therefore, $\pi(2p_{\pi})$ and $\pi(2p_{\pi})$ are also degenerate. Inter-nuclear region Similarly $\pi'(2p_a)$ and $\pi'(2p_a)$ are also degenerate

Thus six molecular orbitals are produced by overlap of two sets of p-orbitals. Three bonding and three antibonding.



Flore of the Copyring data of the control for a



College Completes: Part-I

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Character Bondon

The bond formed by linear (head on) overlap is called σ -bond. While the bond formed by sideways overlap is called x- bond

Since three bonding orbitals are produced, therefore, porbitals can form three bonds. one sigma and two x- bonds.

delative energies (1908)

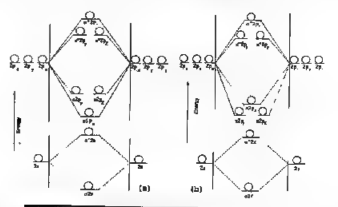
The energies of molecular orbitals are determined by spectroscopy,

The energy of MOs of diatomic molecules such as O1, F4 and their positive and negative ions is in the following order

 $\sigma(1s) < \sigma^*(1s) < \sigma(2s) < \sigma^*(2s) < \sigma(2p_n) < \pi(2p_n) = \pi(2p_n) < \pi^*(2p_n) = \pi^*(2p_n) < \sigma^*2p_n$ л 2p, and 12p, are of some energy. These are called degenerated orbitals.

The energy of M.Os. of distormic molecules such as N_z Be₂ ,C₂ etc. is in the following order

 $\sigma(1s) < \sigma^*(1s) < \sigma(2s) < \sigma^*(2s) < \sigma^*(2s) < \sigma(2p_n) = \pi(2p_n) < \sigma(2p_n) < \pi^*(2p_n) = \pi^*(2p_n) < \sigma^*2p_n$



Side above 0.0 of the hope is a function of the form of the field from $H_{\rm c}$

In $N_{x_0}B_x$ and C_{x_0} is 2 p_y is binding in energy than $\sqrt{2p_x} = \sqrt{2p_{y_0}}$. Where

In case of N_2 , B_2 and C_1 , σ $2p_n$ is higher in energy than π $2p_n = \pi$ $2p_n$. It is due to mixing of 2s and 2p atomic orbitals.

The energy difference between 2s and 2n orbitals is small. Therefore, these are mixed together (hubindization of AO). Thus or 2s and of 2s do not have pure s - character

Chambers, Port-



guillarly, or 2p, and of 2p, MOs do not have pure p-character All these four MOs gain spguillens. As a result, energy of these orbitals changes, of 2s and of 2s becomes more stable the section of the control while of Sp. and o 22, becomes less stable and are raised in anemay.

Since a 2n = a 2p, are not involved in milking, therefore, their energy remains same. Thus energy of a 2p, is talsed so much that it goes higher than x 2p,= x 2p, bonding objials.

This mixing does not occur with O. F. due to large energy difference between 2s and 20. orbitals.

Energy differences between 2s and 2p orbitals are given below. These have been determined by spectroscopy.

Boron = 554 kJ/mal

Carbon = 846 kJ/moi Nitrogen = 1195 kJ/moi

Oxygen = 1595 kd/mor

Fluorine = 2078 kI/mos

BOND ORDER

The number of bonds formed between two atoms by overlap of atomic orbitals is called the bond order.

ip MOT, the band order is defined as

The half of the difference between the number of bonding electrons and unti-handing

It is given by the formula

Bond Order = No. of electron in bunding MO No. of electron in antibonding MO

Generally, valence orbitals are considered in band order culculations.

EXAMPLES

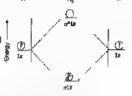
THST PERIOD ELEMINIS

forwation of H, majecule

Electronic configuration of H is 18

MO diagram of H₂ is shown in the fig. Hz molecule has two electrons both in (ofs) orbital and forms a single covalent bond as shown in fig.

Electronic configuration of H₂ is <u>or Isl</u>



Epper After alarmiala person by

Bond order = $\frac{2-0}{2} = 1$

Thus, there is a single coverent between two hydrogen atoms in H, molecule



Colonial Control State

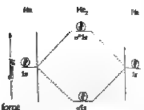
Commence House Com

- Electronic configuration of He la

 Jrle 1s³
- In MO diagram is shown in the lig.
- (4e) configuration is u(1s)² < or4(1s)²
- Bond Oules $=\frac{2-2}{2}=0$

Thus there is no bond Le; no attractive force between two He atoms.

Hence Haltum molecule (He₂) does not exist.



 ≸ quer décise danset, tals protiere di He Leupothéricale

SI COSD PLINOU LELMENTS

Language of S Malerale

- Electronic configuration of N is _nN 1s¹ 2s² 2p_n¹ 2p_p¹ 2p_n¹
- Electronic configuration of N₂ is

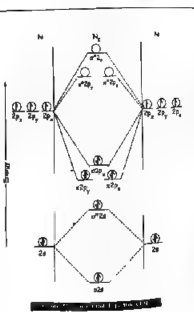
$$\begin{split} & \sigma(1s)^3 < \sigma^n(1s)^3 < \sigma(2s)^2 < \sigma^n(2s)^2 \\ & < \pi \left(2p_p\right)^2 = \pi \left(2p_p\right)^2 < \sigma \left(2p_h\right)^2 \end{split}$$

 Since all the six electrons enter into three outerment bonding orbitals while no electrons enter into antibonding orbitals. Thus

Bond Order =
$$\frac{6-0}{2}$$
 = 3

Thus there is present triple bond between two nitrogen atoms in N_a molecule. i.e. N=N. One bond is o-bond while two others are x-bonds.

Bond Energy of Na = 941 kJ, mol



Bedronic configuration of O is

,0 ls 2s 2p, 2p, 2p, 2p,

. Electronic configuration of O₂ moteonic is

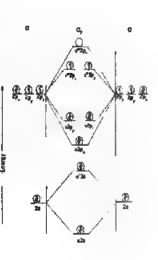
 $\begin{array}{l} _{\sigma(1s)^{2}<\sigma^{*}(1s)^{2}<\sigma(2s)^{2}<\sigma^{4}(2s)^{2}<\sigma(2p_{s})^{2}} \\ <_{\pi}(2p_{r})^{2}=_{\pi}(2p_{s})^{2}<_{\pi}^{*}(2p_{r})^{1}=_{\pi}^{*}(2p_{s})^{1} \end{array}$

 $\mathbf{good\ Order} = \frac{6-2}{2} = 2$

Thus, $O_{\mathbf{z}}$ molecule has a double and between two oxygen atoms i.e; $O\!=\!O$

- O_z has two unpaired electrons in π*2p, and π*2p, orbitals. Due to these in unpaired electrons O_z shows in paramagnetic behaviour.
- The paramagnetic behaviour of Ω₂ cannot be explained by Valence Bond. Theory it is the main success of MOT. Equid O₃ is attracted towards the magnet.

Bond energy of Op = 494 Kill mad



In \mathbb{Q}_2^{-2} , two electrons are added, thus its paramagnetism finishes. Similarly in \mathbb{Q}_1^{-2} two imported electrons are removed, thus its paramagnetism olso finishes. Bond order of \mathbb{Q}_2^{-2} is one and \mathbb{Q}_2^{-2} is three.



Charles Pag-I





BOND FIERGY BOND LENGTH AND DIPOLE MOMENT

Benil Energy Bond Enthalps >

The merops energy required to break one mole of similar bonds in a substan Head bound controlls.

_mi(\$

It is expressed in kJ/mot

- It is the energy required to break an Avogodro's number of bonds. Same amount of energy is reseased when an Auogadro's number of bonds is formed.
- Experimentally bond energies are determined by calorimetry or spectroscopy.

 The enthalpy change in splitting a molecule into its component claims is called enthalpy of atomization.

Jame Character and Board Energy

Bond energy is a measure of strength of bond. Strength of bonds depend upon

- Electronogrativity difference of bonded atoms
- Sizes of atoms
- Bond length

						1 - Carterill	
Bornd	Mond energy thinnel "I	Hund	Bond server (kdpml)*t	Bond .	Front (Bornt America	€ Bejnd émist()' (k5me('''
2.3	348	н-н	436	0-0	145	\$I-H	323
6-6	614	H-F	567	0=0	495	SI-Si	226
C=C	839	H-CI	431	O-H	463	SI-C	301
CH	413	i-i-iir	266	0-F	190	SI-O	368
C-M	293	H-I	299	0-a	203	F-H	155
C=N	615	N-N	163	ы	234	G-F	253
CWN	891	N=N	418	5-6	266	a-a	242
C-0	354	Nen	441	S=5	416	Br-F	237
C=0	759	24-15	391	5-0	623	Br-CI	215
C=O	1072	N-O	201	S-H	339	Br-Br	193
CP	465	N-F	272	5-8	327	Fa	208

A tolar condent band is stronger than a non-polar consilent bond. Consider the greater strength of HCI, due to its polaritonic) character

The H - H bond energy is 436 kJ/mol

.e. . H + H → H_L ΔH = -436 kl/mol

Thus 6.02 × 10 ^{sh} bonds have energy = 436 kJ

MM 1 bond will have energy = $\frac{1}{6.02 \times 10^{33}} = 72.42 \times 10^{-63} \text{ kJ}$

Since each hydrogen atom contribute equally, therefore, contribution of each hydrogen will be 36.21×10^{-15} kJ,

- Similarly Cl Cl bond energy is 240 kl/mox and for one bond it is 39.86×10^{-18} kJ. Thus each Cliatom will contribute 19 93 x 10-23 kJ.
- Hence In case of H = Cl, bond energy must be 36.28 × 10⁻¹³ +19 93 × 10⁻²⁵ = 56.31 × 10⁻²⁵ kd/molecute
- > For one mote it will be 338.9 kd/mol. But actually it is 432 kd/mol. This increase in bond energy is due to polar character of H - Cl bond.

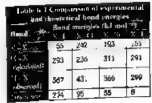
Bond energies generally decrease with decrease in electronogativity difference of the bonded atoms.

Example: In $H=F,\,H=Cl$, H=Br and H=I, greatest E.N. difference is in H=F bond Le. $2.1,\,$ while in H=I, the E.N. difference is the least i.e. 0.4.

Hence, H - F has greatest bond energy (568 KJ/mol) while that of HI is the least (299

- alectronogativities determined by noting difference between experimental and calculated bond energies. e.g. in case of HIX, this difference is largest for HP and least for HI. It shows that HF is more ionic than Hi
- The order of lante character to

HF > HCl > HBr > HI









- - - w_{ij} . The bond amorphis of $C \cong C$, C = C and C = C are to the order

$$C=C>C=C>C-C$$

Thus train bend is stronger than double bund which is stronger than single boar. However, board energy of triple bond is not thrice of single bond. Similarly but energy of double bond is not double then single bond. Thus it shows fast sign bond is stronger than pi bond.

on the group, due to business in bond length for Bond energy decreases do particular type of bond.

BOND LLNGHT

The manufic distance between the muclei of two atoms forming a consists bad a collect bond length

r mit

In SI system bond lengths are expressed in part (1pm = 10^{-18} m).

The bond lengths are measured by physical methods, e.g. electron diffraction, [48] diffraction or spectral studies.

Due to vibratory motion of atoms, bond lengths are not constant. Thus measures has length is actually the average value.

The covalent bond length of a particular bond is almost independent of the natural molecules. e.g. in most stephatic hydrocabons C - C bond length is 154 pm. Sans box length is present in diamond as well.

A baron Superal Carallera Rudi " (1907) b

One half of the distance between nuclei of two similar bonded atoms is a

Constant route of different elements are additive in acture.

Exemple

Covalent radius of Ct is 99 pm, which is half of Ct - Ct bond length Le.198 pm. Similarity

Covalent radius of C is 77 pm. which is half of C - C bond length i.e 154 pm.





Then C \sim CI bond length must be 99 + 77 \approx 376 pm.

(Lawrence, in contain cames, this role does not apply.

Excepte: By adding covalent radii of Si (117 pm) and F (64 pm), the espected bond leigh of Si - F in SiF₄ is 181 pm but actual value is 154 159 pm.

This deviation is due to difference in electromagnitivity of the bonded atoms.

Generally, greater an electronegativity difference, shorter the bond

That calculated values are always higher than actual pakers due to electronegativity digrams. It is because, ionic character is produced in the bond. The potes attract each other and bond length becomes shorter.

Table 6.5 Some selected bond lengths along with in bridication of central atom					
. Сопрения	Hybridization	Bond wit	Band longth Ipm		
Heisan Hillighte	sp² .	B-P	120		
IMAMI Calabatha	spi [®]	B-Cl	175		
Salt Swing.	olij.	Bi H	148		
Silleon (einalleisrale)	कृषे	94 - F	155		
CH (Library)	tg ²	C-C	1 154		
CH.	MD ₂	c-c	133		
CII CHERNON	ağı	c=c	120		
Security Con	\$P ⁸	¢=0	122		

Schallen Sup of Heridization with Bond (small)

In general bonds are shortened by increasing a character of the hybrid orbitals. It is because s-orbital has smallest mean radius. nh (om)

e.g. Bond	Hubridization.	% s - Charms	
C-Cin ethyne	5D	50%	120
C-Cin ethene	₽D ²	33.33%	133
C-C in ethane	40°	25%	154







15.6



s – bonding also reduces the inter-nuclear bond distance

Generally, triple bond is shorter than a double bond, which, In turn is shorter than einche bond

e.g. Bond	Bend length Lorn
C-C	154
C = C	133
C = C	120

From Sin Periodic Told

Along Period

Generally, band length degreeses from left to right in a periodic table due to decrease b atomic size and increase in nuclear charge.

e.g. bond length of C - C is greater than $N \cdot N$ bond length

In a Group

Generally, bond length increases down the work due to increase in atomic size

s.g. Si = Si bond length is more than C + C bond length.

P - P bond length is more than N - N bond length.

h is because due to increase in alcunic radius, effective nuclear charge decreases or electrons. Thus bond length increases.

DIPOLE MOMENT

"The product of the magnitude of one of the charges (positive or negative) and is most between their centers is called dipole mament"

The bond between two atoms of different electronegativities is polar i.e. have positive and negative poles. This produces polarity in the mosecule.

The separation of charges on bonded atoms is called a dipote. It is measured in terms of dipole moment defined as

Let if 'q' is the magnitude of one of the charges and 't' is the distance between them, then

The dipole moment is a vector quantity. It is represented by an arrow, directed form electropositive end to the electronegative end of dipole.

For policitomic molecules, the net dipole moment is the resultant of vector addition of individual bond moments.



Thus from eq.,1) and (2)

$$\mu = \frac{1.6022 \times 10^{-19}}{3.336 \times 10^{-30}} \approx 4.80$$

or $ID = 3.336 \times 10^{-60} \text{ mC}$ (2)

Si unit of choose moment is mC (meter coloumb)

 $10 = 10^{-10}$ est. cm.

W?LICATIONS

Divole Marrierus and Millegelia Stracture

Dipole moment measurements help to determine the shape of molecules.

 $\mu = \{1.6022 \times 10^{-19} \text{C}) \times \left(100 \times 10^{-68} \, \text{m}\right) = 1.6022 \times 10^{-69} \, \text{mC} \subseteq (1)$ mC is a bigger unit Generally, smaller unit Debye (D) is used.

Distance malecules

Diatomic molecules are always linear eg. CO has a linear structure and its dipole moment is 0.12 D.

Thus for a unit negetive charge at a distance of 100 pm from a unit positive charge, we

H H H-184D

Irlatomic Molecules

Both CO_2 and CS_3 has zero dipote moment. Thus these have linear 0 151D

shuthere, in which one dipole cancels the

affect of other

Dispole moment of SO₂ and H₂O are 1.64 D and 1.84 D respectively. It shows that SO₂ and H₂O have angular structure because one dipole does and cancel the effect of other.

<u>letoromia Mpleoules</u>

the state of the s from ents do not cancel the effect of each other it is because NMs has been

has Pyramidal Structure

Patinionic Molecules Both CCl, and CH, have no dipole moment which suggests a regular tetrahedral structure for













Secretory lines Chairmen

Experimental dipole moments can be used to determine the Sage tonic character of bond

The formula is

Tage ionic character =
$$\frac{\mu_{\text{obs}}}{\mu_{\text{test}}} \times 100$$

Where p_{ole} = Experimental dipole resonant

 μ_{balk} =Dipole moment calculated on the bests of 100% charge esperation:

Example 1.

The observed dipole meanant of HF to 1.900. The distance between the charges is 0.917 x 10⁻¹²m. Find the percentage ionic character of the bond.

Solution

Unit positive charge = 1.6022 X 10¹⁹ C

Charge $= q = 1.6022 \times 10^{-19} \text{ C}$

Bond length = $r = 0.917 \times 10^{-10} \text{m}$

$$\mu_{obs}=1.90\,D$$

$$\mu_{\rm max}=2$$

% ionic character= ?

 $\mu_{\text{trains}} = q \times r = 1.6022 \times 10^{-10} \text{ C} \times 0.917 \times 10^{-10} \text{ m}.$

$$= 1.469 \times 10^{-29} \text{mC}$$

Since 1 D = 3.336× 10⁻⁴⁴ mC

Therefore
$$\mu_{inste} = \frac{1.469 \times 10^{-28}}{3.336 \times 10^{-30}} = 4.4 \, D$$

Thus % localic character =
$$\frac{1.90}{4.4} \times 100 = 43.2 \%$$

Hence, 43% of HE bond is long in nature and 57% is complete. The word is makely tralevo



Comprised	Dig do		
	the mental		
H ₂	0.00		
HCL	1.03		
HER	0.78		
HI	0.38		
H ₂ O	.85		
HS	2.85 0.55		
NH.	1.49		
SO	1 63		
H.S. NH, SO, CO, CO	1.49 1.63 0.00		
CO	0.12		
NO ×	0.16		
H _e O _e	2.20 0.00		
CH,	0.00		
CH _a F CH _a CI	1.81		
CHIC	1.45		





THE IS OF BONDING ON THE PHYSICAL PROPERTIES OF COMPOUNDS The properties of substances depend upon the types of bonding present in them.

OTHER DE отс Сопровода

legic compounds are soluble in water but insoluble in non-equipme solvents

- if is because polar water molecules detach cations and among from the crystal lattice by gerbrostatic attractions. Trust, tonic compounds are distolved by <u>hudration</u> of lone.
- . The energy released during hydration is used to break the lettice.
- . Only those crustals are dissolved in water for which, enemy produced during hydration s greater than lattice energy, so that it can break the intice.
- . Many tonic crystals do not dissolve in water because attraction of water molecules cannot overcome the attraction between jons.
- ionic compounds are not dissolved in non-noise solvents like between and hereine.

Cetalent Compounds

- · Generally covalent compounds dissolve in non-poler organic solvents e.g. beneene, effier etc.
- In this case attraction of solvent molecules with the compounds break the intermolecular
- Most covarient compounds are insoluble in H_RO. Few compounds are dissolved by ludregen bonding.

Picetional Nature and Isomerisa

laste Compounds

limic compounds do not show isomerism because tonic bands are non-raid and <u>ion-directional,</u>

Coglent compounds

١

Covalent compounds show isometism because covarent bonds are rigid and directional

P.G. C. H.O shows structural isomerism

... OFOI

Compounds having same molecular formula but different structures and properties are called somers and the phenomenon is called somerism



45.0

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Ass. Charles Suid

desector Kinesies

Jank Compounds Reactions of sonic compounds are servi fest

It is because ionic compounds exist as ions in eq. solution. Chemical reaction there was occur repidly.

e.g. By adding AgNO₃ solution to NaCl solution, white ppt of AgCl are produced a

in this reaction, both AgNO, and NeCl are in longest form. Therefore no bond is to be broken. Only few bonds are formed. Thus reaction occurs rapidly

Condent Compounds

Reactions of covident compounds are generally slow. Because no electrical force as present in them to speed up the reaction and energy is required to break bonds and and new bonds. The molecules undergo a chamical charige as a whole.

Covalent compounds show many types of reactions which depends upon the ways reaction and kind of reaction.

DE IL CHVI, AND SHORT ANSWER QUESTIONS IT VERCEST. Salect the correct statement.

(i) Antionic compound A* B is most likely to be formed when solve to

- (a) The ionization energy of A is high and electron affinity of B is low
- (b) The lonization energy of A is low and electron affinity of B is high
- (c) Both the ionization energy of A and electron affinity of B are high.
- (d) Both the ionization energy of A and electron affinity of B are low

(ii) The member of bonds in nitrogen molecules is

- (a) one or and one a
- (b) one or and two w
- (c) three sigma only
- (d) two or and one is

Sandjuich Reard, 2010, 2012) (D.G. Khun Board, 2011, 2012) 2011 A (Sargosko Board, 2012) (Lahora board, 2014)

(Bi) Which of the following statements is out correct regarding bonding melecular orbitule.

- (a) Bonding motecular orbitals possess less energy than atomic orbitals form which they are formed
- (b) Bonding molecular orbitals have low electron density between the two nuclei.
- (c) Every electron in the bonding molecular orbitals contributes to the attraction between atoms.
- (d) Bonding molecular orbitals are formed when the electron waves undergo constructive interference.

(b) Which of the following molecules has zero dipole moment?

(a) NH₄

(b) CHCI,

(c) H₂O

(d) BF₃

(Nation Board, 2011) (D.G. Khan Board, 2012) (Outcounds board, 2014)

ple character (v) Which of the hydrogen buildes has the highest percentage of it

(a) HF

(b) HBr

(c) HCl

(d) HI

[O/ CLEAN | Colored Bourd 2007] (D.G. Khan Bourd 2007, 2012) [Fluoriphical Bourd, 2009] [Fabriched Scord, 2009, 2013] [Fluoriphical Bourd, 2009] [Fluoriphical Bourd, 2009, 2014] [Colored Bourd, 2012, 2014] [Colored Bourd, 2012, 2014] [Colored Bourd, 2012, 2014] [Colored Bourd, 2012, 2014]

(vi) Which of the following molecules has expeired electrons in unifoculting molecular orbitals

 $^{(a)}O_{a}^{b}$

(c) B₂

(d) F₀

(b) N_k-1 Calculate Bound, 2006) (Protectional Bound, 2009) (Palaken Baund, 2009) (Balcaraline Bound, 2009), 2011) (Palaken Bound, 2009) (Balcaraline Bound, 2009), 2013) (Palaken Bound, 2012) (Calculaten Bound, 2012) (Calculaten Bound, 2012)







College Chemistry, Parisi

of Aus, the

ANSWERS TO MULTIPLE CHOICE QUESTIONS tii: Aus (b)

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No molecule. there are times bonds between two nitrogen atoms one bond a sigma and other bond is pi-bond.

Character Box

to loose electron coolly and electron affinity of ecceptor stom must be high to accept electro (iii) Ans. (la)

Tunic bond is formed by complete transfer of

ordization energy of donor atom must be low

electron from one alors to another Hence,

According to MOL when two atomic orbitals overlap with such other they form two molecular orbitals. Bonding MO and antiborating MO in bonding MO, electron density, is mexicuum between the nuclei while in antibonding MO, electron density is minimum between the nuclei.

herefore, [b] is correct

tel Ans: lab

The polarity of a bond depends upon the difference of electronegativity between the two HF atectronegativity atoms. Since in difference is greatest, therefore, bond between H and F is highly poler it has higher dipole moment of all. Thus, it has highest % ionic moment of all.

NH4, H2O and CHO, ere all unsyntmetrical molecules. Their individues band moments do not cancel the effect of each other, therefore all these have dipole moment

However, BF₃ has symmetrical structure in structure is triangular planer. In this structure individual bond moments cancel the effect of each other Hence, BF, has sero dipole moneys

(vi) Aus. (b) In B_a and F_a molecules all the electrons are paired. In O_2 , two unpaired electrons are present to antiborating MO. However, when these two electrons are removed O_i^{1+} is produced which has no unpaired electrons, in N_{χ} no unpaired electrons are recent twayer, when two electrons are added to it, Nation is produced which has two unpained electrons in antibonding molecular orbitals just like O_s, For MO diagrams see Q126 per end Section 5.4.6

Pill to the blanks Q2.

- (i) The tendency of atoms to attain maximum of eight electrons in the valence stell is called
- (ii) The geometrical shape of SICI, and PCIs can be explained on the basis of ____ hybridization
- (III) The VSEPR theory stands (or
- _ then π 2p, orbital. (iv) For Ne molecule the energy of or 2px orbital is _
- (v) The paramagnetic property of O₂ is well explained on the basis of MO theory of terms of the presence of _____ electrons in two MO orbitals
- vi) The bond order of N₂ is ____ while that of Nez is
- (vii)The value of dipote moment for CS₂ is while for SO₂ is

(ii) eight, webst ride till solt till, valence shell electron pelt republish (iv) greater (v) tillpaired (vi three, zero (vii) zero 1 bl

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86.



- Classify the etatements as tree of false. Explain the res-
- The core of an atom is the atom valence shell.
- (ii) The motecules of nitrogen N a N and acetytene HC = HC are not isoelectronic.
- (iii) There are four coordinate coverent bonds in NH₄* ion.
- (b) A or -bond is attronger than a n-bond and electrons of or bond are more diffused than π - bond
- [v] The bond energy of hetero diatomic molecules increases with the decrease in the electronegativities of the bonded atoms.
- With Increase in bond order bond length decreases and bond strength increases
- (vil) The first foruzation energies of the elements rise steadtly with increasing atomic number from top to bottom is a group.
- (viii) A double bond is stronger than single bond and a triple bond is weaker than a double bond.
- The bonds formed between the elements having electronagativity difference more than 17 are always covalent in nature
- (k) The repulsive force between the two bonding pairs is less than that between the two lone pains
- (xi) The number of covalent bonds an atom can form is related to the number of anpaired electrons it has.
- (xill). The rules which govern the filling of electrons into the atomic orbitals also govern filling of electrons into the molecular orbitals.



Q4. What is a observiced band? Discuss the formation of tools and exculent bonds

Page 324 (Ionic Bond) and Page 326 (Casalent Bond)

Q5, (a)Define lantacition energy and electron affinity. How these quantities change with en increase in the mose number?

Page 319 Nonization energy, and Page 321 (electron affinity)

(b) Explain what do you understand by the term electronogations? Discuss its perhatons in the periodic table. How does it affect the bond strength?

Page 323





College Characters: Party

-57

Q6. Writes the Lands structures for the following compounds (e) H₂SO₄ (to) HUN AIF, (vf) H_iPO, (H) CCI, (HI)CS, (A) HCN (x) Ag(NH_a)₂NO₃ HO,HN (at)

(viiii) K₂Cr₂O₃ $(O_{\mu}H(m_{\nu}))$ (m) ស្ពេចស្ពីស្ពេច (m) (ទី១០ នេទី) (n H:C##N:

Н :Ё: (м) н:Ё s : Ф; н (м) н:Ё s : Ф; н Н :Ё: (м) н:Ё s : Ф; н (м) н:Ё s : Ф; н Н :Ё:

Q7. (a) Explain qualitatively the valence bond theory. How does it differ from malecula

Page 323 (For Valence Bond Theory)

Molecular Orbital Henry Valence Bond Theory

In VBT only valence electrons are involved

in VBT, the two combining atoms do not loose their identity

It does not excisin the paramagnetic behaviour of molecules e.g. O₂ molecule

He₂ molecule

In MOT all electrons of combining atoms at involved in bond formation

in MOT, the two atoms lose their identity

In explains the paramagnetic behaviour

It give idea about bond order. So we can determine whether the bond is single, doubts or triple or even in fraction

Il explains the non-existence of Hez molecular.

the following molecules can be explained with respect to

CI₂, O₃, N₃, HF, H₂\$

stop of Cl₂ molécule

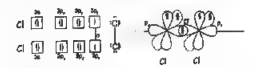
Cls molecule is formed by two chlorine atoms.

The electronic configuration of CL is

17Cl 1s*2s*2p*3s*3p,*3p,*3p,*

Two Cl atoms have half-filled 3p, orbitals. These orbitals overlap to give Cl, molecule will a single bond .o - bond) between two CI atoms.

Diagrammatically, formation of Cla molecule can be represented as



HF motecule is formed by combination of hydrogen and fluorine atoms

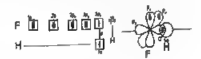
The electronic configurations of F & H are

F 1s22s4 2p, 42p, 42p, 4

3H 141

According to V.B.T. half-filled 2p, orbital of F atom overlage with half-filled 1s orbital of H to form a or-bond and thus HF molecule is formed.

Diagrammatically, formation of HP molecule can be represented as



formation of O₂ molecule

 $\Omega_{\rm I}$ molecule is formed by combination of two oxygen alarm

The electronic configuration of O #

(O 1 nº 2 nº 2 p. * 2 p. * 2 p. * 2 p. *



Two O atoms have half-filled 2p, & 2p, orbitals 2p, orbitals of two O atoms overlap to give a x bond on to give a x - bond, while two 2p, orbitals of two O atoms overlap to give a x bond on to give a x - bond between the head on to give a σ -bond, where is one σ -bond, and one π -bond between two $a_{ij}a_{ij}$ part in O₁ molecule.

Diagrammetically, formation of \boldsymbol{O}_{k} motionals can be represented as



a of N. moli

N₂ molecule is formed by combination of two nitrogen atoms.

The electronic configuration of N is

N 1st 2st 2p, 2p, 2p, 2p,

Two N atoms have half-filled 2px, 2p, & 2p, orbitals. 2px orbitals of two N ato overtap hasad on to give a o - board, while two 2p, orbitals and two 2p, orbitals of two 8 atoms overlap to give two x – bond by parallel overlapping.

Thus, there is one σ_{-} band and two π_{-} bands between two nitrogen atoms in \mathbb{N}_{i} molecule.

Diagrammatically, formation of N₂ molecule can be represented as



Formation of H.S.

Page 339

Q8. Explain VSEPR theory. Discuss the structures of CH₄, NH₄ and H₂O with reference to this therm. to this theory and hybridization schemes?

Page 331 (VSEPR theory, and Page 340, 341 (hybridisation of CH₄, NH₂ and H₂O)

(4) The molecules of NF, and BF, all have molecular formula of the type XF, But they have different structural formulas. Keeping in view VSEPS theory sketch the chape of each molecule and explain the origin of difference in shapes.

In NFs, nitrogen is sp⁸-hybridized. There are four electron pairs present around mitrogen atom. It forms three covalent bonds with three fluorine atoms. there is one ione pair present on nilrogen atom. Hence according to VSEPR theory, it will have a triangular pyramidal structure like NH₈. However, bond angle in NF₈ is 102°, which is less than that of NH₉ i.e., 107.5° It is because F is anger electronegative than N. Therefore, polarity of N. F bond pulls the ione pair of Niclosex to the mucleus. Thus, it exerts greater repulsions on bond pain, hence bond angle s reduced to 102°

In BFz, B is sp² hybridized. It is surrounded by three electron pals. It forms three covalent bonds with three fluorine atoms. There is no lone pair on boron. Three electron pairs are present at a maximum angle of 120° from each other in order to have minimum repulsion between them and form trigonal planar geometry. Thus BF₃ molecule is trigonal planar with bond angle of 120°



Q10.The species NH₂, NH₂, NH₄* have bond angles of 105°, 107.5° and 109.5° respectively, durity these values by drawing their structures.

In NH_{q}^{+} , nitrogen is sp^{3} -hybridized. It forms three covalent bonds and one co-ordinate covalent bond with hydrogen atoms. There is no lone pair perent on nitrogen atom. Hence, according to VSEPR theory, it will have a regular tetrahedral structure with bond angle of 109.5°



In NH, nitrogen is sp³-hybridized. It forces three covalent bonds with hidrogen atoms. There is also present a lone pair present on nitrogen atoms. There is also present a lone pair present on nitrogen atom. Hence according to VSEPR theory, it will have a triangular primaridal. Pramidal structure. Since lone pair exerts more repulsions on bond pairs, literate the bond angle will be reduced from 109.5° to 107.5°



In NH₂, nitrogen is sp⁸-hybridized it forms two covalent bonds with hydrogen atoms. There are also present two ione pairs or phrogen atom.





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tion of the

Harror: accepting to VSEPR theory, it will have a heart structure. Since two lone pair each accept adjustment on board pairs. Hence, the board angle will be exhaud from 109.5° to 109.

QES to Emphase attends original imbeldimentum with reference to sp², sp² and up modes of hyperdimentum for PH₂, C_pH₄ and C_pH₂. Discuss governments of CCI_p, PCI_p and K_p.

We have discussed of constant atoms.

Page 339

The linear guarantey of BoCl₂ reggress that control (to atom is sy-hybridized, u_{to} the linear guarantey of BoCl₂ reggress subset the atoms bonded to it as agree of hybridized v. a control atom realization of hybridized v. a control atom and hybridized at the commerc of (b) at adultational triangle, (d) a regular tetrahedron and

An equietexi trangle - sp² hybridizacion A regular tetrahedron: - 45° hybridisation A triangular bipyramid – dap⁵ hybridization

Q12, tal. Gloss the banks of the unlocaler orbitals theory and discuss the solecule arbitals configurations of the following melecular.

Page 348 (0 Hez

Page 349 alocaler orbital theory explain the paramagnetic character of $0_p \, 0_l^{\gamma}$ (a) O₂

(b) How does malecular and O₂* malecular?

Formation Of O, Molecula.

Electronic configuration of O is O 14 24 2p, 2p, 2p, 2p,

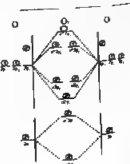
Thus, there are stoken electrons in

O_x molecule.

 O_{μ} has two tenpaired electrons in $\pi^{\alpha}2p_{\mu}$ and x*2p, orbitals. Due to these unpaired electrons O₂ shows paramagnetic behaviour

Electronic configuration of O₂ molecule is $\sigma(1s)^4 < \sigma^*(1s)^4 < \sigma \cdot (2s)^4 < \sigma^*(2s)^4 <$ $\sigma(2p_s)^2 < \pi_s(2p_s)^2 = \pi_s(2p_s)^2 < \pi^2_s(2p_s)^2 = \pi^2_s(2p_s)^2$ Liquid Or is attracted towards the magnet.

Band Order = $\frac{6-2}{2} = 2$

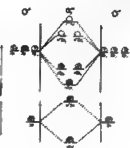


when of $O_{g}^{-\alpha}$ form.)s ${O_{\kappa}}^{*S}$ ion, two electrons are κ first day fourteen electrons in O_2^{-1} loss.

Or has no unpaired electrons in any odd parto this corporated electrons O2** shows danguetic behaviour i.e., it a not attracted and magnet.

Bond Order =
$$\frac{6-0}{2}$$
 = 3



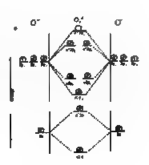


fermation of O_2^{-1} Lan.

In O₂ sion, two electrons are added. Thus, there are Shakern electrons in O_2^{-4} ion.

 $\mathbf{Q}_{\mathbf{z}^{\mathbf{d}}}$ for no unpaired electrons in any citial. Due to this $O_{\mathbf{s}}^{-\mathbf{s}}$ shows diamagnetic thatour, i.e., it is not attracted towards Dgagt,

Bond Order =
$$\frac{6-4}{2}$$
 = 1



(e) Sketch the moleculer orbital pictures of

(42) a (2p_) and a (2p_)

Page 345

(b) O₂, O₂2+, O₂2-

Q 12

(c) He_s and Ne_s

Page 348, MO diagram of Ne is had the $\mathcal{O}_{\mathbb{R}}$ Only electronic candiguration is different.

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July.

(b) Sketch the hybrid urbitals of the speaker, PCl_{j} , δP_{0} , δICl_{j} and NM_{i} .

in PCI, P is ap hybridized. P has four up hybrid orbitals overlap with a critical of three chiertre atoms to form three covalent bonds. One bybrid orbital of P contains lone pair. Thus, PCI, has a trigonal pyramidal gramatry



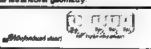
ppP(hybridized state) - Jy - F - r 40° 40° 40° 40°

In SF_{\bullet} , S is $ep^{\bullet}d^{\bullet}$ hybridized. S has six $ep^{\bullet}d^{\bullet}$ hybrid orbitals. its hybrid orbitals overlap with p-probleds of six flour ne atoms to form six covalent bonds. Thus, SP, has an octohedral





in SiCl₄, Si is ep⁸-hybridized. Si has four ep⁸-hybrid orbitals its hybrid orbitals overlap with p-orbitals of four chionne atoms to form four covalent bonds. Thus, SiCl_d has letrahedral geometry.



in NPL,*, N is sp² hybridized. N has four sp³ hybrid orbitals. Its hybrid orbitals evenlap with s-orbitals of four hydrogen atoms to form three sevalent bonds and one co-ordiente covalent bond. Thus NHI,* has tetrahedral geometry



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Annie Mander (EET L'ANNIE) (EET LEET

014. (a) Define hand energy. Explain the serious parameters which determine strongth.

o Hour do you compare the bond strengths of

(i) Polar and Non-polar malecules

Non-polar molecules are formed either between similar atoms (e.g., He) or have animetrical structure (e.g., CCl₄) in H₂, the bond is non-polar while in CCl₂, individual conducted polar Generally, polar bonds are stronger than non-polar bonds.

Polar molecules have unsymmetrical structure. They always have polar bonds and polar bonds are stronger than non-polar bonds.

m wand whonds

Segma Honor	Py Dorod
It to forested when two atomic orbinals graphic with their suces in a consight line.	It is formed when two p-orbitals overlap with their axes pseulid to each other
The averlap of morals orbitals is more than prisond. Hence, a large arrown of energy a released and the bond is strong.	The overlap of montic orbitals is only diaght. Hence this bond is quite weak.
The probability of finding electron is maximum around the line joining the two needs:	The probability of finding electron is manicomin above and between the line joining the two nucles.
A ligner, bond is formed between the atoms, which are not already bonded.	A Pi bond is formed between the atoms, which are already bonded by a signal bond.
Signa band is stronger than pi bond. Example for Cl ₃ , there is signal band between two otherine stores. For Cl – Cl	P) bond is weather thus nigons bond Example. in O ₂ , there are two bonds between two astygen atomic, one is rigous and other is pl. v.e. O = O

Calculate the band energy of H-Br. The band energy of H-H is 486 kJmol' and that of Br.Br to 198 kJmof

The H - H bond energy is 436 killmol

Le.

Thus

6.02 × 10 ** bonds have energy = 436 kJ

1 bond will have energy = $\frac{436}{6.02 \times 10^{23}} = 72.43 \times 10^{-24} \text{ kJ}$ 436





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Since each hydrogen atom contribute equally, therefore, contribution of ϵ_{id} hydrogen will be $-36.21\times 10^{-19}~{\rm kd}.$

Similarly, Br = Br bond energy is 193 kJ/mol and for one bond it is 32.06×10^{-25} kJ.

Thus, each Briatom will contribute 16.03 × 10-44 kJ.

Hence in case o H Br, bond energy must be

 $36.28 \times 10^{-20} + 16.03 \times 10^{-20} = 52.31 \times 10^{-20}$ kl/molecule

Thus, for one mole it will be $52.31\times 10^{-21}\times\ 6.02\times 10^{28}$

= 324.90 kJ/mol,

Actual bond energy of HB: is 366 kJ/mol. This increase in bond energy, due to polar character of $H-Br\,bond$.

Q15. (a) Define dipole moment. How does it help to find out the chapes of molecules?

See Section 6.5.4

(b) The band length of H-Br is 1.4 X 10⁻⁹m. its observed dipole moment is 0.79D/Pm the percentage lands character of the band.

Unit positive charge = $1.6022 \times 10^{19} \text{ C}$ and $1 \text{ D} = 3.336 \times 10^{49} \text{ mC}$

Bond length =
$$r = 1.4 \times 10^{-10} \text{ m}$$

$$p_{\rm obs}=0.79~{\rm D}$$

 $\mu_{\text{tents}}=?$

% ionic character=?

$$\mu_{\text{toric}} = q \times r = 1.6022 \times 10^{-14} \text{ C} \times 1.4 \times 10^{-16} \text{m}$$

$$= 2.243 \times 10^{-29} \text{mC}$$

Since 1 D = 3:336× 10-10 mC

Therefore

$$\mu_{lank} = \frac{2.243 \times 10^{-29}}{3.336 \times 10^{-30}} = 6.723D$$

Thus % ionic character =
$$\frac{0.79}{6.723} \times 100 = 11.75$$
 %



Characteric Parts

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Chartesi Sentine

Q16. PF₁ is a polar molecular with dipole moment 1.02 D and thus the P.F bond to polar. St, being in proximity of P in the Pariodic table, it is expected that St.F band would also be polar but StF, has no dipole moment.

PF, have a triangular pyramidal structure.

This structure is unsymmetrical. Thus in this structure individual bond moments do not cancel the effect of each other. Hence, it shows dipole moment.



SIF₄ molecule has a regular tetrahedral structure. This structure is appropriated. Thus in this structure individual bond moments cancel the effect of each other. Hence, SIF₄ does not show dipole moment.



Q17. Which of the following molecules will be polar or non-polar, sketch the etructures and justify your answer:

(i) CCL

In CCl₄, all C-Cl bonds are poter CCl₄ motecule has a regular tetahedral structure. This structure is symmetrical, it has uniform deciron density distribution. In this structure, individual bond moments cancel the effect of each other. Therefore, its dipole moment is zero releace, in CCl₄, all C-Cl bonds are polar but molecule is overall non-polar.



III SO,

In SO_{L} , all S=O bonds are polar. SO_{3} molecule has a Ingonal planar structure. This structure is symmetrical. It has uniform distribution. In this structure, individual bond thoments cancel the effect of each other. Therefore, its dipole moment is zero. Hence, in SO_{3} , all S=O bonds are polar but holecule is overall non-polar.

(its) MP,

In NF₈, all N-F bonds are polar. NF₁ molecule has a trigonal planar structure. This structure is not symmetrical in this structure, individual bond moments do not cancer the effect of each other. Therefore, it has dipole moment. Hence, in NF₃, all N-F bonds are polar and molecule is also polar.







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(hv) SO₂

In SO_{2n} both S=0 bonds are polar, SO_{2n} molecule has an angular structure. This structure is not symmetrical, in this structure, individual bond maments do not cancel the effect of each other. Therefore, it has dispose moment. Hence, in SO₁, both S=O bonds are polar and molecule is also polar.

Q18. Explote the following:

(I) Bond distance is the compression distance between hen stome. (Guiramente Board, 2009: Sargadha Board, 2010: D.G. Khan Board, 2011, 2012: Guiramenta Board, 2011; See Section 6.1.2

in many course, the distinction between a co-ordinate conclust bond and a cooled band analabas after band formation in NH₄*, H₂O* and CH₂NH₄*. (Labore Bond 201)

A co-ordinate constant band is formed when both the shared electrons are desired

After board foundation in manty cases no distinction remains between co-ordinate covaint and and covalent bond.

in H₆O° ion, experimentally it has been found that all bonds are equivalent. Thus, their is no distinction between co-ordinate covolent bond and covalent bond-

tion of MH₄" for to

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Paration of CH,NH,* Ion to

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mijeths bond angles of HaO and NH, are not 109.5" like that of CH., Although O- and Natoms are op hybridized.

CH₄ molecule has no lone pair and its structure is tetrahedral with bond angles 109.5° However, in NHs one lone pair is present which exerts repulsions on bond pairs thus the book-angle is raduced to 107.5° in case of H₂G, two lone pairs are present which exert none repulsion on bond pairs and therefore, bond angle is reduced to 104.5"

(a) about are more diffused than c-bands.

Johne Scard, 2028, 2012, 2014. Bohrinsipur Bound, 2009. Mukan Bound, 2007, 2009, 2011. Surgadha
 Sand, 2009. Falicikabad Bound, 2007, 2010. D.G. Khen Bound, 2010. Gujraninala Bound, 2010. 2012, 2014.
 Od. Khan Bound, 2012. Rescripted Bound, 2012. Rescripted Bound, 2007, 2013.

a-bond is formed by head on overlap of atomic orbitals of atoms. Therefore, electron density is largely present between the mucles. However, whond is formed by sideways overlap of atomic orbitals. Therefore, electron density is largely present above and below the Hence n-bond is more diffused than o-bond

(i) The absormality of bond length and bond strength in HI is less principant than that of HCL [Lahore Board, 2007; D.G. Khen Board, 2011 Faiselahad Board, 2012 Researched Board, 2012]

in H-Cl, the electronegativity, difference of H and Cl is 0.9.

While in H-1, the electronegetivity difference of H and I is less than 0.9

Dise is greater electronegativity difference in H-Cl, the partial charges on H and Cl in HCl are greater electronegativity disterence at 1995, he and Cl have more attractions and HCl are greater than those on H and I in Hil. As a result, H and Cl habeutour shoremalities hence band length and bond energy shows greater unexpected behaviour ,abnormalities) than that of HI



(cf) The dipole measures of CO_2 and CO_3 are zero, but that of SO_2 to $1.61\,D$ (Gupranuela Board, 2009: Multon Board, 2010: 2/13)

CO₂ and Cb₂ have linear structures which are symmetrical. In these structures included bond moments cancel the effect of each other Hence, dipole moments becomes an include an experience of angular structures in which included bond moments do not be a significant of the decode moments. cannot the effect of each other Triesalone II has dipole morners.

tate The meleting and healing paints of electroneless compounds (i.e. lonks are early high as compared with those of excelent compounds. Why?

CSR Host of improvements and host of additionations. | Buttons Board, 2012, Singarbia board, 2014,

he some comprounds some me hald together by strong electrostates forom of attractors (to of energy a sequent to tune these forces. Hence m.p. and h.p. of routs compounds m Apply (Mark

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he considered companied intermediacións forces use present in the form of dipole-dipte forces and it forming als. These forces are easy to break their ionac forces, blance cover compounds have low in p. and h.p. as compare to fortic orimpounds.

HELLOI Mr. Question herei

Multiple Choice Questions from PAST PAPERS

- 1. Call has some character (Latine sound 2014)
 - (a) 60%
- (b) \$9%
- (d) 100%

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(d) V-inspet

- The geometery of ethere in chousemake west, 2011)
- of Tetrahedical (b) Trigonal stories (c) finear Which inclasses has zero dipole moment? (Guara
 - 40 BF.
- (b) CHC ((c) HOs (a) PAHs 4. The number of electrons shared in \$F₄ molecule is sofa famouri, 2000
 - (b) 12 (c) 6 (m) 4
- 5. The amount of energy released by absorbing electron in the training electron
 - (a) vonization energy
- (b) electron allienty
- (c) electronegotivity
- (d) alcortesion energy
- Formation of chemical bond takes places when datum hand, 1916.
 - bedroeds at various (e)
 - forces of repulsion overcome forces of attraction
 - (c) forces of attraction are equal to forces of reputation
 - (d) forces of attraction overcome forces of sepuleous
 - Carbon dioxide and methone have dipole moment value: (the
 - (a) sero and 1.95 D
- (b) 1.70 D and 1.80 D
- (c) both have seco
- (d) none of these
- 6. Noble grees have murphourn stability and least stability because abuse (a) they are very safe (b) their valence shells are complete
- (d) they are present in zero group
- (c) they are goes 9. Octob rule is not followed in the formation of: Wahanager Stand, 2010; Patenthed Barri, 2011.

 Germania based, 2400, 2010.
 - (a) NF
- (b) CF₄
 - (c) OCL
- de harry (. 1911). 10 long and co-ordinate covalent bonds are present in: 10-per QH (b)
- (a) SO₂

- (b) NH₄C £ (c) C₂H₂ 1). The highest electronogrative element in the periodic table in (Guyannata bount, 2013)

- (a) onlygen (b) nitrogen (c) chionine 12. According to VSERP sheety, the shape of PH₃ molecule is: (a) Trigonal Pyramidal (b) Tetragonal (c) Linear (d) Trigonal Planer



		al Wiss	column to catalog	an Squard, 501 ()							
13.1	Mg becomes Iso ele	scronic with Neon	The second second second	trons							
Į	(a) Looses one electron (b) Looses (wo electrons										
- 1	(c) Gains one duct										
	The radius of No.2	ion is prompted to	pard, \$915)	(d) 95 pm							
	Feel E	(b) 93 pm	(c) 94 pm	fed so hu							
15.	Bond order of $\mathcal{O}_1^{2^*}$	is @maipled board,		40.75							
	(a) 2ero	(b) One	(c) Two	(d) Three							
16.	The dipole momen	nt of SO ₂ is. (Make	Board, 2013)	(1) 0.00							
	44 0.0D	њ 0.12 D	(c) 0.95 D	(d) 0.78 D							
17.	Which one of the it	followings has the l	highest bond orde	er? (Melson Board, 2013)							
	(a) O_2^{1+}	(b) O ₃ *2	(c) O ₂ ⁻¹	(d) O ₂ -2							
18	The most stable ele	ements are: (Lebera	Board, 3009								
	(a) Halogens	(b) Lithium family	y (c) Noble g	gasses (d) None of these							
	The hybridization			2009)							
	(a) sp	(b) sp ¹	(c) sp ³	(d) Not Hybridized							
20.	The geometry of a	there is: (********	f Board, 2009)								
	(a) Tetrahedra I	(b) Trigonal Plane	er (c) Uno	ear (d) V-shaped							
21	The value of dipol	le moment of CS ₂ i	is Bulancipus Board	d, 2009)							
	(a) 0.12D	(b) Zero	(c) 1.61D	(d) 0.95D							
22.	200P. 2013)		2	ed on the basis of (Surgodia Souri,							
				eary (d) CF theory							
23	The bond order of			podha Board, 2009)							
	(a) zero	(b) 1	(c) 2	(d) 3							
24.	The four equivalen	nt sp ^a bybrid orbita	uls in space are at	an angle of 10.6. When Board, 1908							
	(a) 120°C	(b) 107.5°C	(c) 104.5°C	(d) 103/2/C							
25.	. Bond angle between										
	(a) 104.5°	(ъ) 107.5°	(c) 92°	(d) 95°							
26.	. S.I unit of dipole r			7 (b. 12							
	(a) pm	(b) Debye	(c) mc	(d) all							
27	Which pair has trip										
_	(a) CO ₂ , SO ₂	(b) NH ₂ , PH ₃	(c) BF ₃ , AlH ₃	(d) H ₂ O , C ₂ H ₂							
28	The decrease in a	tomic radius is sma	all when we travel	from left to righ in transition							
	elements, Scal to 2	Zn _{io} , Y _{ae} to Cd _{et} du	12 to: (Felesiahad R	Sourd, 2010)							
	(a) valence electro	ons (Ł	b) intervening elec	etrons							

Callent Characters Parts 100 (c) Shielding effect (d) screening effect 19. Which of the following compound has a coordinate covalent bond? (San (a) NH₄Cl (b) NaCt (c) HCl (d) ALCI 30. The carbon atom in C₂H₄ is (6-gods, Seed, 2010) (a) sp⁹ hybridized (b) sp² hybridized (c) up hybridized (d) dap! hybridians 31 MOT was proposed by (Resolvink Souré, 2010, D.G. Khan Bouré, 2011) (a) Morley (b) Werner (c) Kossel (d) Mulikan's 32. Which of the following molecule has a coordinate covalent bond. (2.6). 86m Board, 2010; a) NH₃ (b) AICI, (c) HCI (d) NH₄CI 33. Total number of sigma bonds in ethyne (CH = CH) are (D.O. Khan Sourd, 2010) (a) five (b) three (c) two (d) four 34. The SI unit of dipole moment is (Latina Board, 2011) (a) doubte (b) Debye (c) Coulometer (d) Nam-2 35. Total number of bonds in C₂H₄ mojecule ask (Lobora Board, 2021) (a) six (b) Four (c) Five-(d) Eight 36: The most electronegative element is (Faholabor Board, 2011) (b) F₂ (c) O₂ $\langle d \rangle H_a$ (a) N₂ 37 Dipole moment of CO₂ is: (Labore Souré, 2007) (d) 2.2 D (a) 1.84 D (b) Zero D (c) 0.95 D 38...in sp² bybridization, the orbitals are oriented at an angle of: (Labore Board, 2007) (b) 120° (c) 180° (d) 90° (a) 109.5° 39. The geometry of BeCl₂ is (Felesished Board, 2007) (a) Linear (b) Planar trigonal (c) Tetrahedral (d) None of these 40. The mojecule which cannot for co-ordinate covalent bond with H* ion is (c) PH_a (d) CH₄ (a) NH_a (b) H_aO 41. VSEPR theory was proposed by (Sergodin Road, 2014)
(a) Kossel (b) Lewis (c) Nylhotm and Gillespie (d) Sidwick 42. The shielding effect is responsible for: (a) The decrease in nuclear attractive influence over the valence electrons (b) The increase in nuclear attractive influence over the valence electrons

(c) The decrease repulsion between Nucleus and Inner Electrons
(d) The increase in attraction between Nucleus and Inner Electrons

The state of the state of

THE BUTTON



Answers to Multiple Choice Questions from Past Pupers.									
Q#	-	Q#	Ans	Q#	Ans	Q#	Ans	Q#	Ans
1	(c)	2	(6)	3	(d)	4	(c)	5	(6)
6	(d)	7	(c)	8	(b,	9	(d)	10	(b)
11	(d)	12	[a,	15	(b)	14	(d)	15	(b)
16	lai	17	(b)	18	(c)	19	(b)	20	(b)
21	(b)	22	(c)	23	(d)	24	(d)	25	(c)
26	(c)	27	(c)	28	(b)	29	(a)	30	(6)
31	(d)	32	[d]	33	(b)	34	(c)	35	(a)
36	(b)	87	(6)	38	(b,	39	(n)	40	(d)
41	(c)	42	(a)					1.	4

Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK-I

SHORT & LONG QUESTIONS FROM PAST PAPERS

OCTET RULE

- (I) Explain Octet rute
- (2) What is actet rule? Why certain elements do not aboy it? (Bahawalpar Board, 9010)
- (3) what is ociet rule? Give two examples of compounds which deviate it. (Midian Board, 1911: Lahora Board, 2907, 2012; Midian Board, 2012)
 (4) Write the Lewis structure of NH_aOH and H_aSO₄ (Midian Board, 2012)
- (5) Write the Lewis structures of CS₂ and CCl₄ (Seegothe Board, 2014)

ATOMIC RADIUS, IONIC RADIUS

Shart Questions

- (1) Define atomic radius. How does it vary in groups and periods? (Laters Board, 2009)
- (2) Why atomic radius cannot be measured accurately (or precisely)? (Sempothe Board, 1910) werfo Roard, 2018, 2010
- (3) Why atomic radius increases down the group? (Revelpted Board, 2007; Sagodia Board, 2009) n Bound, 2010)
- (4) Why ionic radius of a cotion (positive ion) is smaller than its parent atom? (Labora Bosto) soos OR The radius of cation is smaller than atomic radius but that of an arion is greater than atomic radius of an arion is greater than atomic radius. (Nation Speed, 2011; Labora Boord, 2010, 2014)
- (5) Anionic radius is always lorger than parent atom. Why? (Surgosine Board, 2007) OR With the lorder radius of Cl. Inc. Inc. of the Control of the Contr ionic radius of Cl 1 lon increases from 99 pm to 181 pm? (Lakers Board, 2018) of

Grand Charliston, Ports

374

A. M.C. Transfer

Compare the radius of chloride ion with its parent atom. Give reasons, there

- 8008 6) What is interioric distance? Explain with example, (D.S. Khee Board, \$610)
- (i) Write short note on atomic and land radii. Give examples

ONIZATION ENERGY

- Sen Greetens

 (i) Define ionization energy. What factors influence the tonization energy/potentia? As
- (2) Whi the ionization energies values decrease down the group although necular charge increases. (Fulsolational Beard, 2008: D.G. Many Board, 2006. Behavelyan Board, 2009, 2011: D.S. Then Beard, 2009, 1012: Surgodia Board, 2012: Outrements Board, 2015) of Group VIA ster.

 [3] Why forticalion energy increases from left to right in periods? (D.G. Khao Based, 2007)
- Intedi Board, 2009)
- (Supermond Standardson Energy) potential. Hous does it vary for trend in periodic table?

 (Supermole Stand, 2005, Multon Secret, 2009, 2011, 2012; Sergothe Sowd, 2010; Februidest Sowd, 2017, 2012, 2013; D.G. Rhen Board, 2011, 2012; Behavedper Board, 2012; Labors Sound, 2013, 2014)
- (b) Define ionization energy. Why second ionization energy/potential of an element is always grader than first tonisation energy/potential? (Labor Seed, 2008; Saloumber Seed, 1968; Saloumber Seed, 2010)
- (6) Jet landation energy of Mg is lower than its 2nd landation energy. Explain (Remission)
- [7] Explain that ionization energy is the index of metallic character. (D.S. Riem Board, 2009: Segunda Board, 2013)
- [ii] Orbital nature also affect the tonisation energy. Exprein. (0.0. Rum Board, 2010)
- (1) Define ionization energy. Give factors upon which it depends. How does it vary in periodic table? (Resolpted Board 2009: Matter Board, 2015)

ELECTRON AFFINITY

Short Overtions

- [1] Define electron affinity. Discuss its trent in periodic lable. (And Reshab Board, 2012) OR How do electron affinity values vary in the periodic table? (Mater Boord, 2010)
- (a) Why the first electron offinity for most of the elments is negative while the second electron offinity for all the elements is positive. (D.G. Ki Board, 1908; (Labore Board, 1908). Sergedhe Board, 2009; DE Why the second electron affinity of oxygen is positive? (Reference Board, 2009)
- (a) What factors influence the electron offinity? (Foliational Board, 2011)
- (ii) Define electron affinity. Name the factors affecting it. How does it vary in the periodic table? (Lahere Board, 2013)

ELECTRONEGATIVITY

- Short Questions

 The County of the County of
- (2) What is electron scativity (b) electron affinity (same source, and the least electron scation along the least electron scatter a





Colon Continue Cont

- Define electromagnishy. How do electromagnishy unless very in the periodic tols. Sealing from the Seas, 1912; Felenters Burret, 1919; Palenters Seas, 1912; Malles Seas, 1917; Palenters Seas, 1917; Malles Seas, 1917; Malle (3) Define mischaneg), take and the found, 1912; have a second in particular table, grandely, described in particular table, grandely, a second for Distance electromagnitisty. Give its trend in particular table, grandely a
- avery
 this rice electromagnitude values of aliensaria help us in uncircularizing the nature of
 sharmfall bond? president theret date date flour does the electromagnitude difference date.
- From Oversion $(1)^{-1}$ What is electromagnetistly? Discuss its curiation in the periodic table. How does it offers to
- (2) Define the term electronogenisty. Discuss its certains in the periodic table, the

SOME SCHOOL CONNESSY BOND, COOKENATE COVALENT SOME

- Why the carry ori kondi ere directional? endan inc
- (2) Separate the Solicions projection in polar or non-polar status (6) CO₂ and CH₂OH (6) CO₂ for H²
 (7) No board in chaminty is 100% ionic Justify it. Granded Basel
- Adres Sept. 2007)

- 16 Define tonic bond. One one economic thank tentes from 2010;

 15. What is non-polar condent bond? Give in comple. (miles form), 2001;

 16. Define polar condent bond with example. (fundami flore), 2001; forgotte file.
- (7) Tall the number of alexinon poins shared in CH₂Cl and 1 '21 many bonds are pair.
- Michaelber Stand, 2003.

 She Daffine coordinate considere based and give its harr accomples obspressive found 500 flampeds flame, 5000 Francisco Stand, 5011 Flampeds Flame, 5011 Flampeds Flame, 5011 Flampeds Flame, 5011 Flampeds Flame, 5010 (Or English the Internation of Lessin model 500, flamped Flame, 5010)

 The Continue the Internation of Lessin model 500, flamped Flame, 5010)
- Fine a coordinate condition band differs from constant band? (Labora Board, 1917)
- ium constant and co-cordinate constant bonds with two examples in e^{\pm} the Street Street

VALENCE WALL PLECTHON PARK REPLACEON THEORY

- (1) Write down two postulates of VSEPR theory, (Labor Board, 1618: Holes Bo
- (2) Write the order of repulsion between electron pairs according to VSEPR THEFT (Belleman Series Series)
- (2) Why the love pairs of electrons on an atom accupy more space? therein Laters Board, 2019: Sundipolis Sensit, 2011 Sugardia Sensit, 2011 Sundipolis Sensit, 2010; Why repulsive Society are less between lone pairs than board pairs? plants Sundipolis Sensit, 2010;
- (5) What is the basic assumption of VSEPR theory, and discuss the shape of a molecular DEF-a constitution of the shape of a molecular NOTE. white Band, MIR NOTE. (BF g containing three electron pairs? Makers Board, 9919. San BP , molecule is on page

Charleton Paral



- Why the man despition the geometry of Nill, replacible occurring to volumes shall electron pair.
- Mily and No. have different band angles heaty me hand angle is 107.5° while in NF, the hand angle is 102° manager from the an inwh Freed, 2010; Oil in 1971, the

- (i), Give three posturates of VSERP theory. Also discuss the strature of BeCl, in the light of
- (i) Write the postulates of VSEPR theory and discuss the structure of NH, with reference to (Barganilles Barrel, 3974)
- (i) Explain VSEPR theory. Discuss structure of SO, according to the theory, and then bend
- # Discuss with two suitable example VSEPR theory of covalent bond. Judge Same 2007.

 [5] Write the main postulates of VSERP theory and explain the structure of annisms on the
- bosis of this theory (Labora Sund, 2012)
- [6] Write down the expected features of VSEPER theory. Also discuss the cases of water and emmonia molecules, diagnée band, 2007)
- (7) Write the postulates of VSEPR theory and discuss the structure of methans with reference to VSEPR theory, distant Breed, 2007.
- SI Give important postulates of VSEPR theory required front, Mrs. Supplies Ser.
- th Explain VSEPR theory with

MUNCE BOND THEORY, HYBEROGATES

- Short Countings
 (1) Write two points of Volence Bond Theory. (Owner, 1) Write two points of Volence Bond Theory. A Board, 2011)
- (2) Sigms bonds are more stronger than pi band. Why? standard the rd, 2007; Friedshad Baard, 1889: Gapta ule Royal, 2011 Bergeette Board, 2011, 2028, 2010; OR Why pf bond is weaker than sigms bond? Resemble Same 200th
- Ordine orbital hybridization and operapping of orbitals or some in
- With the bond anigets of H₂O and NH₂ are not 109.5° like that of CH₄, although oxygen and nitrogen coorse are egi-hybridized Why? Sugments bond. 2001 Little Bond. 2009. Sphridtection Justify it. (Bequate Bond, 1910)
- (5) Deviation in bond angle in water (H₂O) molecule is more than in ammonia thirt, min Servic 2012)
- molecuse although both have tetrohedral structures. (0-per What type of hybridization takes place in boron trifluoride molecule? Comment on it.

Consider Sures, 2000)

(Consider Sures, 2000)

(Consider Sures, 2000)

(S) Explain valence bond theory (VBT) with examples shouling the overlapping of different



(2) What is orbital hybridization? Explain the structure of CH₀ on the basis of by hybrichestion. (Patentina Sand. 2009; Labor Sand, 2014)

hybridization. Classify it and discuss the structure of ethens (CH₂ = CH₃ _[4] (sybridization) (Philips Board, 2010, 2021, Gajrannia Board, 2011, 2013)

(4) Define attantic orbital hybridization. With its help how we can describe the geometrics of Digina diamac oronal reportationals. It was an extension by bridge tion? Explain of the problem hybridization with two exemples (Guyermin Bend, 2008; D.G. Khm Bond, 1911; Bules & PFS. Sugadio Seast, 2013)

(5) Write a note on sp hybridization with an example of contrient. (Asset Real

(b) Define and classify stomic orbital hybridization. Discuss the structure of BF 2 in the light of orbital hydricitations, (Februaria) Source, 2007)

(7) Define up? hybridization and on its basis exploin the structure of atheres, (Fall

MOLECULAR CHRITAL THEORY

- Short Occurred

 (1) How molecular orbital theory justifies that Helium atoms cannot make the He_i manual. 2004, States Sand, 2008, Supplied Sand, 2009, 2013; Olf How molecular orbital theory explains that helium is monoclomic? Printing Bears, third on Draw diagram of He molecule according to MOT Also give its frond order (Nation Board, 2013) Oli He, molecule is not formed. Why? (thinse Board, 2007)
- (2) Give the sequence of molecular orbitals in N₂ molecule. (the
- | Silver, 2009 | Statement of VET? Administration of the D.S. Rhou Statement Statemen 277 Alaban Saurd, 2009: D.S. Filam Second, 2010: Labore Se offs Second, 2011 Maham Second, 2013)
- (5) Helium is diamagnetic in nature, Justify? (1-hors found, 2011)
- 16. Define bond order Mention bond order of N₂ molecula. (D.G. Khan Board, 1912)
- (7) Define for Exploins hand order. Colculate the bond order of nitrogen for hydrogen of Chrysle, phogodie Mand, 2007; Lebers Stand, 2007; April B. which Marrie 2015 Later Street
- (B) Define bond order and what is bond order of $O_2^{2+\alpha}$ S-sumbod Bond, 2013) (S) Why MOT is suprior both VSEPR and VSEP strength Bond, 2013)
- (10)
- Shetch the molecular orbital picture of O₂, (Gujermeth Board, 2013) When is found order? Flow it can be colculated? (D.G. Khan Burrel, 2007) (110)
- (12)
- Why the energy of antibonding molecular orbital is higher than corresponded Foreign recincular orbitals? chapaths Board, \$807. Labour Board, \$812) (13)
- Differentiate between bonding and antibonding molecular orbitals. (Luture Bond. 200 0.4. Sheet, 2018)
- (1) Explain the molecular orbital structures of following molecules on the basis of the MOT
- (ii) N, Phirogen) (iii) O₂ (copper) dishare liquid, 30.12)

 (ii) N₁ Phirogeni (iii) O₂ (copper) dishare liquid, 30.12)

 (iii) Explain paramagnetic nature of copper on the basis of molecular orbital than hand, 50.00; p. 6. Shap hand, 50.07. Outsimitals Sent, 10.01; Labory Secret, 10.11, 10.16

Office Chambridge Parts

38.3

percribe the bonding in O₂ according to molecular orbital theory and applain its (g) Describe the control of the cont

What is the basis of molecular orbital theory and explain peramagnetic nature of oxygen (the model from 2009)

- SOND LENGTH, HOND ENERGY
 Ser Gentlems
 (I) Why a double band is stronger than single bond? provided bond, 1996.
- [2] Define (a) electronegativity (b) band energy
- Compare band strength of polar and non-polar bonds. (Nambbal Band, 2016; Nambbal Band, 2016; Nambbal
- (4) Define bond length and bond energy, (Felo olahud Bered, 2007, 2012; Seepedin Based, 2012;
- (5) Define bond energy with two parameters which determine its strength. (Co-Long Quantions

Define band energy. Discuss two factors effecting the band energy (Laters Box

- Define bond energy, and explain the various parameters which determine its strength.
- Define boral energy and explain with the help of a suitable example the effect of lank disposter on its value. (Sequelle Board, 2013)

MPOLE MOMENT

- Seri-Questions

 (b) Why the clippole moment of CO_0 is zero and that of H_0O is 1.85D, possibled Bound, 1888.
- Why the dipole moment of CO_2 and CS_2 is zero and that of SO_2 is 1 61D. (Laboratorial)
- (i) Why the dipole moment of CH, is zero? (Males in
- Hough the Roge ionic character of the polar band can be determined? the standard and standard an
- Disole moment of CO is 0.120 but that of CO; is zero Why? granatus
- Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

 Apr. 2013.

 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

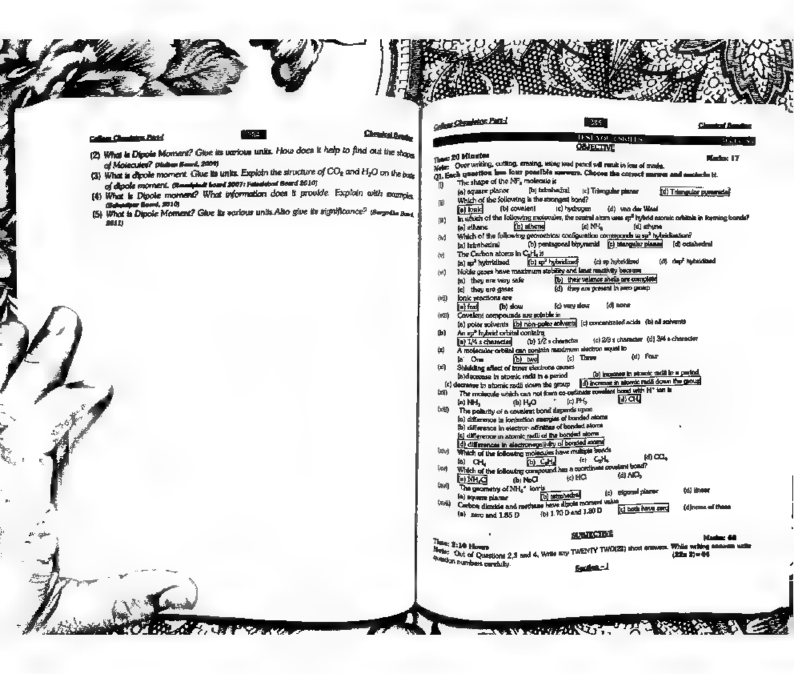
 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

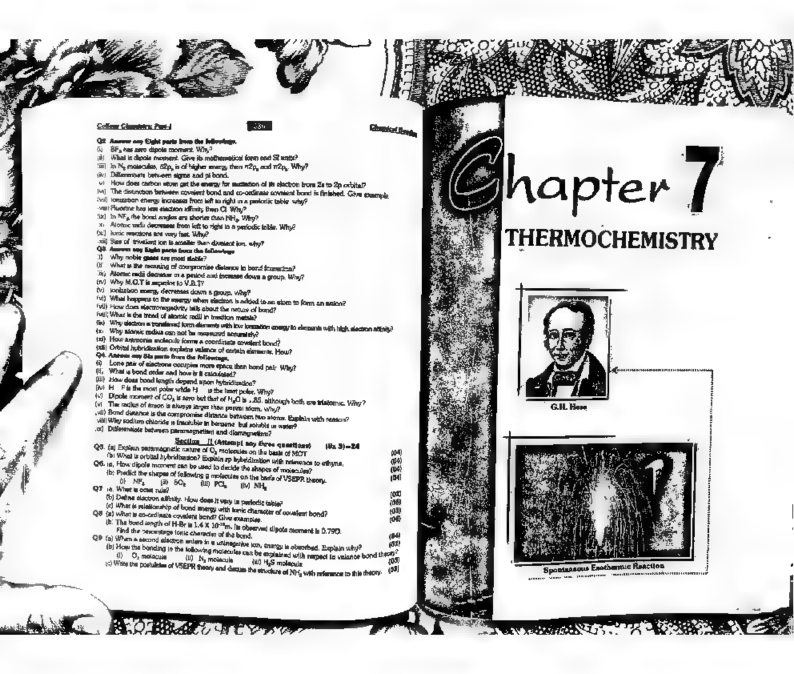
 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.

 Define (or What is) dipole moment. Give its various units for S.I. units, a.c. as 207, 2019.
- Define dipole moment. Give its mathematical expression. (Buselond Seed, 2012. I
- South State Comment. Give an insurance of the state of the state of SO₂ is zero? Access the state of SO₂ is 2.61 D but that of SO₂ is zero? Access there is a state of the state of t
- Why CS_x molecule is finear while SO_x molecule is angular? do.0. Khom Board, 30 (2)
- Why CS, molecule is threat when any more does it explain the geometry of CO, and the dipole moments. Give its SI units. How does it explain the geometry of CO, and the and Bit a molecular and







CONTENTS

Chacker-7

Thermochhemistry

INTRODUCTION SPONTANEOUS AND MON-SPONTANEOUS REACTIONS SYSTEM, SURROUNDING AND STATE FUNCTION

INTERNAL ENERGY AND FIRST LAW OF THEEMSCOPPINAMECS

HEAT CHANGES AT CONSTANT PRESSURE (ENTHALPY)

Processor and American (LAC) Enthalpy of prescribes (LAC) Enthalpy of formation (LAC) Enthalpy of adventuation (LAC) Enthalpy of neutralization (LAC) Enthalpy of combustion (LAC) Enthalpy of solution (LAC) Enthalpy of solution (LAC) Enthalpy of solution (LAC) aurement of anti-pleas

HESE'S LAW OF CONSTANT HEAT SUMMATION The Born-Haber cycle

Objection and short surrectioned (exercise)
himselved problems (exercise
and business MCOs and Short Operations

Other Chambers Pare?

17.5





MESSICHON

The branch of science, which dash with energy changes, to known

Thermochemistry

The branch of chant effect thermochemistry stry which deals with everyy changes in a che

Why heat is evolved or showfood to a chemical resection?

During chemical reaction, bond breaking and making takes place

Bond breaking absorbs energy, and band making orleases it.

in characted reserviors, emergy required to break boards a not could to the energy evolved for bond making. Different substances have <u>different</u> energies and thus <u>total energy of</u> standards a six-way <u>different</u> from the <u>total energy of anothers</u>.

This difference of energy is either evolved or absorbed during a chemical traction. This is

called heat of reaction.

TIPES OF BLACTIONS

On the basis of absorption or evolution of heat, reactions are classified as

(i) Exothermic reactions

(ii) Endothermic reactions.

ZOTH BURE BLACHONS

The practitions in which heat is excited are known

to such reactions, total energy of the products is less than the rocks energy of the reactions.

This difference of energy is avoived as heat.

The heat is given out and paragraphic of the against day above the room temperature.

Also, some time heat is but to the surpossition and temperature fails to room temperature.

Embersolls

The freshed best of reaction is indicated by neuritys size.









350

Emmola:

COper $C_{ee} + O_{bat}$ $\Delta H = -393.7 \text{ kJ/mol}$ (ii) ∆ H= 285.58 kJ/mat Hasi . Va Oasa H_2O_0 (0) N_{max} , $3H_{\text{max}}$ 2NH ... AH= -41.6 kJ/mol

and the second

The reactions in which heat is absorbed are known as audathermic reactions.

The reactions of the energy of the products is greater than the total energy of the such reactions total energy of the products. This difference of energy is absorbed as heat. Thus, heat required for reactions obtained from reactions and thus temperature of the system falls below the monopolitic and formal the system. temperature After some time heat is absorbed from sterrounding and temperature again rise to room temperature

The absorbed heat of reaction is indicated by positive sign.

Emmoles:

 $H_{\text{thin}} + I_{\text{cut}}$ 2HL, ΔH= +52.96 kJ/mor Name + One 2NO. ∆H= +180.51 kJ/mo!

Difference between Exothermic and Endothermic reactions Prochaman Reactions Endothernia Revertous

The reactions in which heat b

absorbed are known as endothernic

In these, total energy of the product is

" these, temperature first fall that is

more than that of reactant.

to room temperature.

The reactions in which heat is evolved are known as excellentate reactions.

In these total energy of the product is was than that of seactant. In these, temperature first rise then fall

to room temperature The heat retensed is indicensed by

negative eign. Example:

+ O20 - CO20

The heat absorbed is indicated by positive sign. Example $N_{2\omega} + O_{3\omega} \rightarrow 2NO_{\omega}$ $\Delta H = +180.51 \text{ kJ/mol}$

reactions.

ΔH= - 393.7 Ы/mo

hagarmax ags (I multitres, a) Therewich airs).

Thermochemistry tells about the energy or heat contents of the system industrial reactions. This knowledge is transfer and an industrial or the system of the system industrial or the system of the s reactions. This knowledge is used to explain chemical contents of the evident the Housever best of reaction can be assumed to explain chemical conding and chemical could not the second flust.

However heat of reaction can be accurately determined only for the reactions. Thus,

Chambers Foots

Ja-

SPONTANEOUS AND NON SPONTANI OUS IMACTIONS Sontaine Q6.

parties to the spontaneous and non-spontaneous processes? Give assumption OF INTANEOUS REACTIONS

A process which takes place on its own without external help and a A process which were process on he was whose scarmer new and the

- , It is a matural real process, therefore, it is unidirectional and preventible process. Emmoles
 - 1: Water flows from higher level to lower level
 - 2 Neutralization of a strong acid with a strong base $NaOH_{pdj} + HCl_{pag} \longrightarrow NaOl_{pdj} + H'_{g}O_{qg}$
- 3. Reaction of Zn with CuSQ, solution

 $Zn_{(a)} + CuSO_{4(a)} \longrightarrow ZnSO_{4(a)} + Cu_{(a)}$

is the reaction, blue colour of CuSO₄ solution disagness due to nedox reaction.

Some spanianeous reaction requires a small input of energy to start. But once they are started they go on by itself until complete

e.g. Burning of coal and hydrocarbons in air.

Coal does not react with Both N_{α} and O_{α} are present in atomosphere. But they do not react. This reaction takes place, when energy is provided by lightning air by itself. This reaction is started by a sperk, when coal starts burning the reaction goe to completion by itself.

***SPONTANEOUS REACTIONS

- A process which does not take place on its ama is celled a non-speni
- I does not occur in nature. It is reverse of montaneous process.
- Sould non-acoutation's biodesics can pe things aboutabloons per computering among of them, from external source. Emmoles

- t. Pumping of water uphili
- Transfer of heat from cold interior part of reingerator to the hot surroundings.

3. Reaction of N_3 with O_1 to form NO

Both N_g and O_g are present in atmosphere. But they do not react. This reaction takes Negl Mr and Or are present to describe the second section is provided by lightning bott.

Region is provided by signatury over the special section and non - positive out the section and intermediate between special and non - positive out







Cellus Charles Part-

3/12

STRUCTION CORESPONDANCING OF REACTION

Experience shows that spontaneous processes occur by the decreese of energy and be Experience shows that spontage to be said that all exothermic reaction to transferred to the surrounding. Thus it can be said that all exothermic reaction to is transverse to use surrounding. The in the state of the spontaneous, However there are many endothermic reactions, which are spontaneous,

$$H_{i}O_{m} \longrightarrow H_{i}O_{in}$$

 $\Delta H = 44 \text{ kJ/mos}$

 NH_4Cl_{ad} $\longrightarrow NH_4^*_{(ad)} + C\Gamma_{(ad)}$ $\Delta H = 15.1$ keVinos Hence energy change is not a criterion for spontaneity of reaction.

Spontaneity of reaction is actually Judged by the free energy change in terms of entropy of system

Difference between Spontaneous and Non-Spontaneous Processes

	Spontoneous Process
	A process which takes place on its own
	without external help and move from
	non-equilibrium state to equilibrium state
	is called a spontaneous or natistal process.
ż	It is a natural real process

It is unidirectional and Imporsible process

Exemple.

(i) Water flows from higher level to lower level (II) NaOH + HCl + NaCl + H2O

A process which does not take glass on its own is called a non-spontaneous

It usually does not occur in nature.

Some non-spontaneous processes car be made apontaneous by continues supply of energy from external source Example

(i) Pumping of water uphill (ii) $N_{244} + O_{245} \rightarrow 2NO_{45}$

SYSTEM, SURROUNDING AND STATE FUNCTION

SYSIUM

Any real or transferary part of the universe that is under study is called

A chemical system is usually a substance undergoing a chemical change

SURROUNDING

Everything that is not part of the system is called surrounding

BOTTVDARY

The real or imaginary surface separating the trounding is called boundary.



39.1

Symples

- in experimental work, a specific amount of one or more substances forms a system e.g. 1 mole of O_a in a cylinder fitted with a piston is a system. The piston, the cylinder and all other objects outside cylinder are surroundings.
- 2. Water contained in a cup is a system and the cup, air around it and the table on which it is present are all surroundings.
- g. If reaction between Zn and CuSO4 solution is taking place in a flask as shown in Rours, the contents of flask is a system while flask and air around it is surrounding.

The condition of a system is called a state

When a process occurs, the state of system is changed

Belonation

Consider a water system at a given T and V. This initial condition of water is called initial to Now if water is heated in a beaker, its condition will be changed. The final condition of hemstem is called final state.

A comparison of final and Initial state talk us about the change taking place in a system. Determine :

Let T₁ and T₂ are the temperatures at initial and final state of the water system then dames in temperature change in state, will be

$$\Delta T = T_x - T_x$$

1-10V 0-8

It is a mairrescopic property of spoten which has definite volume for initial and final was and it is independent of the pash through which change take place.

Conventionally, capital latters are used to represent a state function.

Estimplant Volume (V), Temperature (T). Enthalpy (H), Internal Energy (E) etc.

Let initial volume of gas is V_1 , its volume can be changed to V_2 by changing temperature or pressure. The change in volume is given by

$$\Delta V = V_z - V_z$$

This change can be brought about by changing pressure or temperature.

Shoe V is a state function, therefore, AV is independent of the path through which the tog takes place (i.e by changing T or P)



College Characters: Part-I

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Thermochant

INTERNAL UNITED

it is the sum of all the energies of all the particles (i.e., stom, ions and malecular)

It is denoted by 'E' It depends upon kinetic and potential energy of the system

Le. E = KE + PE

0.00

It depends upon the motion of molecules (i.e., translational, vibrational, rotational),





or a Calculative analysis of

tel Engran:

It is due to binding forces between the particles. It includes inframolecular forces such as covalent bond and intermolecular forces such as van der Waal's forces.

- Internal energy is a state function i.e., it depends only on the initial and final state of a system and notion the path through which change takes place.

 The internal energy of a system cannot be determined. Only changes in internal energy.
- denoted by A.E., can be determined

le, A£=E_e-E_r

LEARCH TRANSALITATION OF SYSTEM AND SPREADINGS WORK AND HEAT

When a system undergoes a change, energy is transferred into or out of the system in $\hbar^{\mu\rho}$ fundamental ways i.e., heat and work,

WOHK

Work is a form of energy and is defined as the product of force and distance.

Work - Force x Distance

It is denoted by w. It is not the property of system, Hence, it is peth function, it is not a etale function.

Sign Conception for a

Work done by the system is measure .e. - w Work done on the system is positive i.e. +w Charlet vi. Parts

395

intra of the

The Shunit of work is Joule (d)

Officer unit is erg

 $1 \text{ erg} = 10^{-7} \text{ J}$

there are many types of work

Chemical mark

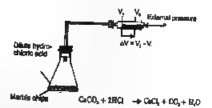
lightenistry most common type of work is pressure-volume work

Consider the reaction between CaCO3 and HCI as shown in fig. The CO2 produced things like voltime form V_1 to V_2 against constant pressure ${}^{\rm P}$ So, the pressure-voltime work

$$W = -P\Delta V$$

where, P = External pressure,

AV= Change in volume



ji 13

it is the quantity of energy that flows across the boundary of a system during a change is state, due to the difference in T between system and surroundings.

it is denoted by q.

his not the property of system. Hence, it is path function, it is not a state function.

im constitution for q.

that cained by the system is positive ..e. +q

lies of by the system is nenetive. i.e. - q

their of q.

The Silver is these for time IS and

Other unit is calonic (cal).

- cal = 4,184 J









College Chambion, Part-I

Let What is the first law of thermodynamics? How does it explain that (i) $q_a = \Delta E(0) q_{a-1}$

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CIRST LAR OF THERMODYNAMICS

The law of conservation of energy is also called the first law of thermodynamics states. Energy can neither be created nor destroyed although it can be changed form one for

OR The energy of the system and surrounding is conserved.

A system cannot destroy or create energy. However, it can exchange energy with is surroundings in the form of heat and work.

Consider a gas enclosed in a cylinder having a piston. If some heat 'q' is added to a sale. & some work 'w is done on it then its internel energy is changed from 'E, to E2' The men change is equal to the sum of both heat & work so that total energy of the system & surroundings remains constant. Hence, the mathematical form of first law of thermodynamics

i.e;
$$\Delta E = E_1 + E_2 = q + w$$
 ____(1)

ΔE = change in internal energy of system.

q = heat added to the system;

w = work done on the system.

Sim Co

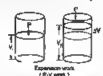
 ΔE is negative when system loses energy $-i.e.-\Delta E$ ΔE is positive when system gains energy

.e; +ΔE Work done on the system is positive i.e. + w

Work done by the system is negative. Let - w

Heat gained by the system is positive i.e; + q

Heat lost by the system is negative $- k \epsilon_i = q$



Processor with manager

TEAT CHANGES AT CONSTANT CORP. OF

At constant valume, let the heat supplied to the system is 'q,' then first let if thermodynamics can be written as

$$\Delta E = q_s + w$$

$$\Delta E = q_a - P \Delta V$$

Since volume is constant, therefore, $\Delta V = 0$

Thus
$$\Delta E = q_a$$

It means that heat exchanged at constant volume is used to change the internal energy the system and no work is done.

Charleton Part

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MALCHANGES AT CONSTANCIPE SSURE A NUCLEPT

graps

Enthology of a system is the sum of its internal energy and product of pressure and

june (PV). It represents the total heat content of a system." Methernatically

$$H = F + PU$$

Explanation

At constant pressure, when heat is supplied to the system, a part of this heat is used to notes the internal energy of a system and a part is used to do P-V work. Thus enthalpy is med to represent the change in state of the system.

Enthalos, is also called total heat content of the system.

iroperties

- 1 It is a state function
- 2 lb unit a <u>doule .Jl</u>
- 3. It cannot be determined for a system in a given state Only change in enthalog can be determined.

Since enthology is

Change in enthalpy is given by

$$\Delta H = \Delta E + \Delta \text{ [PV]}$$

$$\Delta H = \Delta E + V \Delta P + P \Delta V$$

At constant pressure,
$$\Delta P = 0$$

$$\Delta H = \Delta E + P\Delta V_{\perp}(1)$$

According to first law of thermodynamics at constant P

$$\Delta E = q_p - P\Delta V \underline{\hspace{1cm}} (2)$$

Put eq (2) in eq (1)

$$\Delta H = Q_{\bullet} - P\Delta V + P\Delta V$$

$$\Delta H = Q$$

it means that heat exchanged at constant pressure is used to change the enthalpy of a

Most of the chemical reactions are corried out at constant pressure, therefore, A H is more Important than ΔE for a chemist.





Charleton Cartal

Deference better n Internal energy and entiralpy

Income may		Enthalpo
It is the local energy of the system.	1	It is sum of the internal energy and the product of pressure and volume (PV) of the extern
It is the sum of all types of kinetic and potential energies of a system:	2	It is sum of the internal energy and the product of pressure and volume (PV) of the system.
Methaniotically, if is given as $E = KE + PE$	3	Mathematically. It is given as H = E + PV
If it denoted by 'E	4	It is denoted by "H"

Exercise Q 7 (b)

How will you differentiate between ΔE and $\Delta H 2$ is it true that ΔH and ΔE have the same values for the reactions taking place in the solution state

AH is the heat change at constant pressure (Enthalpy Change) and A E is the heat change at constant volume (Internal Energy Change)

Difference between internal energy change and enthalpy change

		_	
	Internal Literge Change 4 (L)		Enthalpy Change (AH)
	The amount of heat evolved or	ш	The amount of hear evolved or absorbed
	absorbed at constant volume is		a constant pressure is called enthalpy
п	called internal energy change	ш	chance pressure is cause entraipy
	(q. = AE)		-
	Mathematically		.q. =ΔH)
1			Mathematically.
	$\Delta E = q + w \text{ or } \Delta E = \Delta H P \Delta V$		$\Delta H = \Delta E + P\Delta V$
4	It is demoted by AE.	3	It is denoted by \Delta !
	-7 -te.	•	It is denoted by \(\frac{1}{2} \)

Relationship Between ALE A. La. Salaton

Change in Enthalpy is given by

AH = AE + PAV

For reactions involving only solids and liquids, or in solution state the change in volume's zero. Le $\Delta V = 0$

Hence

AH = AF

Thus ΔH and ΔF have same values for reactions taking place in solution state.

Peri-I 295 people I har 2 produce of H₀ and 1 mole of O₁ at 100°C and 1 torr pres of general water, 484.5 kd of snarpy are enabled.

(Sorgodia Sound, 2012: Mukin Board, 2013: Bahasaipur Board, 2012) What are (a) AH (b) AE for the production of one mole of H₂O The exaction is The resolved is for two motes of H_0O_{tot} , therefore, heat evolved for one mote at constant P will be $q_p = \Delta H = \frac{-484.5}{2} = \frac{242.25 \text{ kJ/mal}}{1}$ (1) Δ n=No. of moles of product - No. of moles of reactionts $\Delta r = 2 - 3 = -1$ T = 100°C +273 = 373 K P = 1 atm R = 8.314 J mol ⁻¹ K⁻¹ Since $\Delta H = \Delta E + P \Delta V$ $\Delta E = \Delta H P \Delta V$ (2) PV = nRT $\Delta (PV) = \Delta nRT$

or $V\Delta P + P\Delta V = \Delta nRT$

Since P = constant, therefore $\Delta P = 0$

PAV = AnRT

= -1 × 8.314 × 378 = 3100 J

αPΔV = 31kJ

 $\overline{h}_{10}^{\rm b}$ amount is for 2 moles of $H_2 O_{00}$, therefore, for one mole we have

 $PAV = \frac{3.1}{2} = 1.55 \text{ kJ/mol} . _ (3)$

Put value of ΔH from eq (1) and value of $P\Delta V$ from eq (3) in eq (2), we get $\Delta E = \Delta H - P\Delta V$

ΔE = 242.25 -(-1 55) = -240.7 M/mol

when State

The natural physical state of a substance at 25°C (298K) and 1 atm pressure is known thanks of the substance s.p.

The standard state of CO, is gas the standard state of HaO is inquid.

the standard state of Fe is solid.



47/0

Express Q8 (b) Define the following enthaspies and give two examples of each.

- (i) Standard emhalpy of reaction
- (#) Standard enthalpy of combustion
- (th) Standard enthalpy of atomization
- (iv) Standard enthalpy of solution

Enthalps of Reaction (Sec.)

The standard enthalpy change which opening when the number of moles of renewals indicated by the belanced chemical equation react together completely to give prosder stendard conditions.

All the reactants and products are in their standard physical states.

 In exothermic reactions, entirator of product is less than entirator of reactant. Here heal is evolved and AH? is negative, e.g.,

$$\begin{array}{cccc} C_{ac} + O_{bas} & \longrightarrow & CO_{bas} & \Delta H = -393.7 \text{ ki/mot} \\ Z_{M_{abs}} + O_{mas} & \longrightarrow & 2M_{a}O_{cl} & \Delta H = -285.8 \text{ ki/mot} \end{array}$$

In endothermic reactions, entirally of product is greater than anthalou of reactants, Hence, heat is absorbed and AH," is positive

AH=+ 180.51kJ/moi



talah tananahahir

The change of enthalpy when 1 male of the compound is formed from its element, $u^{\pm i}$ standard state is known as standard enthology of formation of the compound.

Examples.

$$Mg_{0i}$$
 + ½ O_{2gi} \longrightarrow MgO_{0i} $\Delta H_i^a = 692 \text{ kJ/mol}$
 C_{0i} + O_{3gi} \longrightarrow CO_{3gi} $\Delta H_i^a = -393.7 \text{ kJ/mol}$

Landadpy of Atmaisation ($\Delta H_{\omega}^{\alpha}$)

The entirelyst change when one mole of gaseous atoms are formed from the situation and are standard conditions is collect standard enthology of etemization of an element. Example

Enthalpies of atomization can be determined by various methods.

Chambers Parket

99.1

Exercise Q9

peps head of neutralization. When a divide solution of a strong acid is neutralized by a divide petit hear of a strong base, the hear of neutralization of a strong acid is neutralized by a dilute skillon of a strong base, the hear of neutralization is found to be nearly the same in all the How do you account for this?

mades of Neutralization (AH2)

the smouth of heat evolved when one mois of hydrogen lone (H*), from an oots, reset applicably with one mote of hydroxide lans (OH), from a base to form one mole of with it called eathulpy of neutralization.

Emploie:

The enthalpy of neutralization of NaOH with HCl is 57.4 kJ/mol

Esthalpy of neutralization of a strong sold with strong base is always approximately constant La. 57,4 kJ/mol

Everyptes

HCI is a strong acid and NaOH is a strong base

Both lonises in water completely

On mixing these solutions, H* and OH* lons react together to form water and heat is owed. While Ne* and Cl. lone are set free in such solution.

Thus, heat released during this process is actually the heat of formation of liquid H₂O rem.H* and OH* ions.

$$Na^*_{[aq]} + OH^*_{(aq)} + H^*_{[aq]} + CI^*_{(aq)} \longrightarrow H_gO + Na^*_{[aq]} + CI^*_{[aq]}$$

$$H^*_{(aq)} + QH^*_{(aq)} \longrightarrow H_2Q_{(q)} \qquad \Delta H_0^* = -57.4 \text{ kJ/mar}$$

Equivalent amounts of all strong acids produce same amount of H* ions in solution. Also individent amounts of all strong bases produce same amount of OH ions in solution.

Since, the net reaction is between H^{\star} and OH^{\dagger} ions, so, the heat of neutralization by strong acid with strong base is always constant [-57 4 ki/mol).

whater of combination (ΔH_{i}^{0})

The mithelpy change when one male of a substance is completely barnt in excess of a substance, a tender standard conditions is valled standard enthalpy of combustion of a substance.

Contract of

 $C_2H_4OH_{10} + 3O_{Rig} \longrightarrow 2CO_{Rig} + 3H_8O_{10} \text{ a H}_2^2 = 1368 \text{ keVmod}$



Cartaday of salatroo(& 🚓 J

The amount of heat absorbed or sectord when one mole of a substance is dissolved in much solvent that further dilution results in no detectable heat change is called only of solution of substance.

Example

- Enthalpy of solution of NH₄Cl is +16.2 kJ/mol. During this heat is absorbed from surrounding and solvent is cooled him andothermic process
- Enthalpy of solution of Na₂CO₂ is -25 ki/mol. During this heat is released and temperature of the solvent rises. It is an exoting

MEASOR MENT OF ENTRACTES

Exothermic and endothermic reactions can be identified by noting the temperature diang. For accurate measurement of AH calorimeters are used.

Two types of calorimeters are generally used

- 1. Glass calorimeter
- 2. Bomb calorimeter

How AM can be determined by using glass Calorimeter? (Faterlahed Board, 2009)

Specific Heat

The amount of best require
increase the T of one gram of
substance by one Relative
it is expressed in J g ⁻¹

1 OASS CALDRAST IN III

THE RESERVE AND ADDRESS.

It consists of an ordinary class, which is insulated from the surroundings by cotton wool as shown in the fig. It is provided with a stirrer and a thermameter for noting T change.



Stoschlometric amounts of substances are placed in

calorimeter. When reaction starts, heat is either evalved or absorbed and thus Tichard The T of the system is noted before and after the chemical reaction. The different

Forms Charles Commercial give the T change (A T) If mass of reactants is "m" and specific heat of reaction mixture is "s" than start hear "q" evolved or absorbed is determined by the eq.

 $Q = m \times 0 \times \Delta T$

College Characterre, Part !

10.5

2 ation of 100 cm² of 0.5M NaOH at 28°C with 100 cm² of 0.5M HCl at 28°C, Appendix hant of water= 4.2 J g f K-1 (Gyranenia Board, 2012, Multar Board, 2012) Solution:

... Volume of acid/ or base used = V = 100 cm³ Initial lamperature $= T_1 = 25 \, ^{4}\text{C}$ Final temperature $= T_s = 28.5 \, {}^{\circ}\text{C}$ Rise in temperature = AT = 28.5 25 = 3.5 °C = 3 K Specific heat of water = 8 = 4.2 3 g 1 K-1

 \Rightarrow Since density of solution (water) $= d = 1 \, g \, cm^{-3}$ Total volume of solution $= V = 100 \, cm^3 + 100 \, cm^3 = 200 \, cm^3$ So, mass of solution $= m = d \times V$ $= 1 \times 200 = 200 \,\mathrm{g}$

 Thus, heat evolved is given by $\begin{array}{rcl} q &=& m\times s\times \Delta T\\ &=& 200\times 4.2\times 3.5 = 2940\ J=2.94\ kJ\\ \text{or} &=& 2.94\ kJ\ (exothermic reaction) \end{array}$

→ Since Molarity of acidy or base used = M = 0.5 M, therefore 1000 cm³ of solution contain HCI = 0.5 moles 100 cm² of solution contain HCl = $\frac{0.5}{1000} \times 100 = 0.05$ moles

0.05 males of HCI react with 0.05 moles of NaOH Thus heat of neutralization is given by 0.05 moles of HCl produce heat = -2.94 kJ = -58.5 8J/moi

Hence heat of neutralization = $\Delta H_0^0 = -38.8 \, \text{kJ/mol}$



Exercise Q8 (b)

How do you measure the heat of combustion of a substance by bomb calcrimster?

зочи съгония так

It is used for accurate determination of heat of combustion of food, fuet and other illibetances.

A STATE OF THE PARTY OF THE PAR



40.4

Construction

College Characters (Sept.)

- It consists of a <u>pulindrical steel vessel</u> I steel bond), lined with gnamed to prevent computers and a screw valve at the top.
- A platinum crucible is present inside steel bomb.
- There is also present an electrical unition coll.
- The bomb is immersed in a known mass of H₂O in an insulated coloringier as shown in fig. The temperature is measured with a thermometer.



- Topics form to share the A known mass (about one gram) of the test <u>substance</u> to placed on a platinum crucible. The lid is closed and
 - cougen is supplied inside until its pressure is 20 atm. The calorimeter is immuned a known mass of water. The initial temperature of water is noted. The substance is the conted electrically by ignition coil
- The water is stimed and its temperature is noted continuously after every 30 sec. To maximum temperature is noted from thermometer.
- The difference of initial and final temperature gives the change in temperature (i.e., AT)
- If number of moles of a substance burnt is 'm' and specific heat of whole system (colorimeter etc.) is 'g' in kel K-1g'', then heat evolved or absorbed 'q is determined by the formula.

The heat capacity of a system is the product of mass and specific heat and is defined in

* The quantity of heat required to alongs the temperature by 1 K.* Thus
$$q = c \times \Delta T$$

Exemple 3

10.16 g of graphite is humit in a bomb calorimeter and the T. rise recorded to \$87.5. addition enthology of combustion of graphite if the specific heat of the calorimeter (book water etc.) is 86.02 to R.

Selution

Mass of graphits = 10.16 gAtomic mass of graphite = 12 g

Moles of graphite = $m = \frac{10.16}{10} = 0.843$ moles 12

Heat capacity = $c = 86.02 \text{ kJ K}^{-1}$ Rise in temperature = $\Delta T = 3.87 \text{ K}$



or Pari-l

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Hest evolved during combustion is given by

0.843 moles of graphite produce heat = 332.89 kJ

= 332.69 =395 kJ/mol 1 mole of graphita produce heat 0.843

- Since reaction is exothermic, therefore, heat of combustion of graphite will be negative. $|a_{-}| \Delta H_c^0 = -395 |\omega|/mol$

Exercise Q11.

(a) Define and explain Hess's top of constant heat summation. Explain a with examples and give the application

(b) Hess's law helps us to calculate the heats of those reactions, which cannot be normally auried out in a laboratory. Exploin it

HESS'S LAW OF CONSTANT HEAT SUMMATION.

G.H. Hess experimentally discovered it in 1840.

In chamical reaction takes place by different ways, the net energy change is some, rejudiest of the route by which the chemical change occurs, provided the initial and final sister are the some

"The net amount of heat evolved or absorbed in a process, including a chemical charge, is the same solution the process takes place in one step or in several steps.

inclunation.

Canader a process occurring as shown in the figure

Mathematically from figure

$$\Delta H_1 + \Delta H_2 + \Delta H_3 = \Delta H_4$$

Thus if system neturns to its initial state in a cyclic

 ΔH_d ANd ΔR,

Process, we can write
$$\sum \Delta H_{(spide)} = 0$$

Hance not enthalpy chance in a closed cycle is aways could to zero. If the initial and final TO SELECT THE PARTY.

Hand's little is simply an application of Law of Conservation of Engree

PROOP.

College Chambric Both

Sedium carbonate can be prepared in two ways

First way

 $2N_{b}OH_{total} + CO_{total} \longrightarrow N_{b}CO_{a,tot} + H_{b}O_{dt} \quad \Delta H_{b} = -89.08 \text{ kJ mot}^{-1}$

Second way

NaOH + CO+ --- NaHCO_x $\Delta H_2 = -48.06 \text{ kJ mol}^{-1}$ $N_BHCO_a+N_BOH \longrightarrow N_{B_a}CO_a+H_BO \quad \Delta\,H_a=-41.02\,\mathrm{kd}\,\,\mathrm{mol}^{-3}$

According to Hear's law

 $\Delta H_1 = \Delta H_1 + \Delta H_2$ 89.08 = 48.06 41.02 -89.08 = -89.08

Hence Hess's law is proved

IMPORTANCE AND APPLICATIONS

There are several processes for which A H cannot be determined directly. These are

- ΔH_1^α of Al_2O_3 and B_2O_4 cannot be determined. It is because, a protective laws is present over the surface of element that resist in complete burning.
- (ii) The compound CCl₄ cannot be prepared directly from <u>graphite</u> and <u>chioring</u>. Moreton CCl₄ is also not easily decomposed into its constituent elements
- (iii) ΔH_i^o of CO cannot be measured directly due to production of CO_2 along with it. For such reaction heat of reaction is determined indirectly by using Hess's law

Example: (Determination of AH, for CO.)

AH," of CO cannot be measured directly due to production of CO, along with it. It also be determined in following ways.

Combustion of carbon can occurs in two ways.

First June

When C is burned in excess of O_{t_0} if is directly converted into $CO_{\tilde{x}}$ $C_{pp} + C_{mp} \longrightarrow CO_{np} \qquad \Delta H_1 = -393.7 \text{ kJ mol}^{-1}$

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Second Way

When C is burned in limited supply of oxygen CO is formed. This CO on further oxidation gives CO.

CO

 $\Delta H_z = ?$

CO₁₀ + % O₁₀₀

COMO

Δ H_a = −283 kJ mol ⁻¹

According to Hess's .aw

$$\Delta H_x = \Delta H_x + \Delta H_0$$

$$\Delta\,H_2=\Delta\,H_1\,-\Delta\,H_3$$

$$\Delta M_{\rm g} = -393.7 - (-283)$$

 $\Delta\,H_0=-110.7$ kJ mol $^{-1}$

Hence heat of formation of CO₆₆ is -110.7 kd moi⁻²



What is lattice energy? How does Born-Hober cycle help to calculate the lattice energy of NoCl?

HIL BORN-HABER CYCLE Latter Energy

The amount of energy released when passous ions of apposite charges, combine to gles one male of a crystelline louic compound."

$$Ne^+_{co} + Cl^+_{co} \rightarrow NeCl_{co}$$

∆H=-787 W mole**

Barn-Heber Cycle

iblebased on the principle that

The sum of energy changes for a closed opolic process is zero, if the teitial and final

Born-Haber cycle is used to determine lattice energy of ionic crystals. Lattice energy contact be determined directly. It is determined by a closed Born-Haber cycle.

brample: (Determination of lattice many of NaCl)

For NaCl, a closed Born-Haber cycle can be drawn as shown in figure. Thus, according to Heas's law,

Na + 1/2 Class 4H + Na 1/2 Class

 $\Delta H_{\text{int}}^{o} = \Delta H_{\text{i}}^{o} - \Delta H_{\text{x}}$ _____(1)

AHT. NO CI'M AH



College Chambring Parts

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Thus if ΔH_1^* and ΔH_n are known, then $\Delta H_{\rm int}^*$ can be calculated

 $\Delta\,H_{\nu}^{\alpha}$ of NeCl can be measured in a calorimeter and its value is = 411 kJ mo) $^{-1}$

$$\Delta H_i^a = -411 \, \text{ke} \, \text{mol}^{-1} \, \underline{\qquad} (2)$$

AH, can be calculated as follow

The overall process is

$$Na_{bp} + V_b Q_b \longrightarrow Na_{bp}^* + Q_{bp}^* = \Delta H_x = ?$$

This process involves following steps

Assembation of Ne_N to Ne_N .

Energy required for this process is determined from values of its heat of fusion, heat of vaporization and specific heat capacity.

$$\Delta H_{e} = 108 \, \text{kJ mol}^{-1}$$

2. Conservior of No. to No. in loss,

This requires contraction energy of New which can be determined by spectroscopy.

$$\Delta r_{\rm k} = 496 \, {\rm kg \, mos^{-1}}$$

Atomisation of Class to Class

Energy required for this process is determined by spectroscopy.

$$^{1/2}Cl_{24} \longrightarrow Cl_{44}$$

$$\Delta H_{\rm el} = 121 \, \rm kJ \, mol^{-1}$$

4. Construion of Clg to Cl* gr

This process is the election affinity of Cl. Its value can also be determined experimently.

lence

$$\Delta H_{a} = \Delta H_{al}(Na) + \Delta H_{b}(Na) + \Delta H_{ac}(Cl_{g}) + \Delta H_{g}(Na)$$

Put values of $\Delta H_{\rm c}$ from eq (3) and $\Delta H_{\rm f}^{\rm c}$ from eq (2) in eq (1) , we get

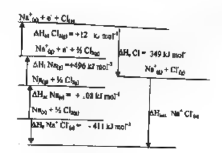
$$\Delta H_{\text{lat}}^{a} = 431 - (376^{\circ}) = 787 \text{ M} \cdot \text{mol}^{-1}$$

Hence lattice energy of NaCl is -787 kJ mol * It represents the forces of attractional NaCl covatals. Lattice energies are used to explain structure, bonding and properties of joint compounds.

The complete Born-Haber cycle for NaCl is shown in the fig.



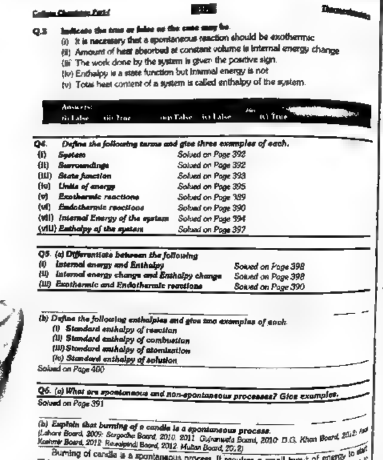
District Contacts



Indicated In Section 1	And the Part of th
	The far character of the strong and and the far the far and and the far and and the far an
to make a now that distribution to the column and produce to now the column and produce to the column are equal to the column	for absorption that a describer, that is approximately found that a describer or the advantage of the found that a section of the advantage of the found that a section of the advantage of the found that a section of the section of
The above or heat many of a classical market of countries of countries of a classical market of countries temperature and persons in spicial countries of subdimention of countries of subdimention to benefit countries. (a) Subdimention of classical countries of classical countries of countr	College of Cliffs Coll
in Example can ratificate be consisted over electroqued. (b) One have all strongs can be increasioned into an equivalent encount of other kinds of the horse an equivalent encount of other kinds of the formation of the changes in independent of its path (Continuous production of the changes work without supplying an equivalent encount of local to possible. (First to global process, the first changes at constant present (q ₀) and at constant entered to global process. (b) q ₀ < q ₀ (c) q ₀ < q ₀ (d) q ₀ = q ₀ /2 (constant and thin) and 1000 (suppose that 101 thin there bound there is constant to the constant of the constan	An a resultationable reservant of an exp sett a boss. Hence there is a second of the s
(a) hear of seastion (b) hear of formation (c) hunt of neutralization (d) hear of combustion (d) hunt of neutralization (d) hunt of combustion (d) hunt of combustion (e) hunt of neutralization (f) hunt of combustion (h) had of combustion (h) hear of season whether it is beenging about to (h) hour or season stage. It is known as (h) hour show (h) hour of conservation of energy had been appeared from a 2013; pherophed heard, 2013.	The substance undergoing a physical or a chemical change forms a chamical (ii) The change in internal energy be measured (iii) Solid which have more than one crystalline forms possess values of heats of formation. (iv) A process is called if it takes place on its own without any outside assistance (v) A is a macroscopic property of a system which is of the path adopted to bring about that change.

10 miles

THE PARTY OF THE P



Burning of candle is a spontaneous process, it requires a small input of energy to set

This reaction is started by a speck When candle starts burning the reaction god completion but had 50 to 10.

completion by faelf. So it is a spontaneous process.

Charles Park 41: it is it true that a non-aprenteneous process happens in the universe? (Resulpted Board, 2010 D.G. Wan Board, 2011, Bahas mansouse process name: happens to the autoporae Justiffy, Englanusia Board, (f) Non-spontaneous processes generally, do not occur in nature by itself. However these (i) Non-sponsor to be carried out by continuous supply of energy e.g. water never flows from processes to higher revel. It is a non-spontaneous process. However this process can be done by using pump and supplying energy. All Some processes may occur in nature under specific conditions, e.g. consider the plowing reaction $N_2 + O_2 \rightarrow 2NO$ Both N₂ and O₂ are present in atmosphere. But they do not react. This reaction takes place, when energy is provided by lightneng boilt (a) What is the first law of thermodynamics? How does it explain that (9) $q_{\nu} = \Delta E$ $(H) \ q_p = \Delta H$ Solved on Page 395 h) Have will you differentiate between AE and AH ? In it true that AH and AE have the stan calues for the reactions taking place in the solution state. Solved on Page 398 98. (d) What is the difference between keyl and temperature? Write a mathematical relationship between these two parameters. Difference between Heat and Temperature Lemperstate ** 11 9217 ****** 70 It is the quantity of energy flows that

Flows across the boundry of system Due to T difference b/w system and

depends, upon the amount of

It is not the property of a given

Hear flows from hot body to cold

It is community determined by

sundunding.

likis a path function

Substance

body

It is the measure of average K.E. of

It does not depend upon the amount of

It is the property of a given system.

h is determined by thermometer

Temperature difference causes the flow

carticles of a substance

of h<u>eal.</u>

6

It is a state function.



College Chemistry, Part-I

414

Mathematical relationship between heat and temperature is

 $q = m \times s \times \Delta T$

q = m x ε x α s.

Where 'q is the amount of heat evolved from a system, when its temperature, changed by AT and a is the specific heat of the system.

(b) How do you measure the heet of combantion of a substance by bomb colorinates) Solved on Page 403

Define herr of neutralization. When a dilute solution of a strong seld insertrolized by a dilute solution of a strong base, the heat of neutralization found to be marry the same in all the cases. How do you account for this?

Solved on Page 401

Q10. (a) State the lase of thermochemistry and show hose are they based on the fu law of therecolynamics

That Lace of the conschemistra:

The heat of formation of a compound is always equal to its heat of decomp with opposite eign.

Example Heat of formation of CO₂ is

Heat of decomposition CD₂ is

$$CO_{kig}$$
 \rightarrow $C_{kij} + O_{kig}$ $\Delta H = +393.7 \text{ kJ mol}^{-4}$

Them, energy is not created not destroyed. Hence, it is based upon the first lat of thermodynamics

Second Law of Thermorteaustre Alless's Law.

The net amount of heat evolved or absorbed in a process, including a cherito change, to the same whether the process takes place in one step or in separal steps Mathematically from fig.

$$\Delta H_1 + \Delta H_2 + \Delta H_3 = \Delta H_4$$
Thus if system setums to its initial state in a cyclic ΔH_1
 $\Delta H_2 = 0$
 $\Delta H_{2} = 0$

$$\Sigma \Delta F_{loyald} = 0$$

Thus, energy is not created nor destroyed, irlence, it is based upon first lar thermodynamics

Charles Charles Port-I

415

respections of equations? Often three examples, What information do they (Mulan Board, 2013: Labore Board, 2012, 2014) Wilhest to a the tion do they

pometr Multan Board, 2013: Labore Board, 2012, 2014)
The thermochemical equations are the chemical equations in which physical glasses of the actants/products and amount of heat evolved or aboscreed is given.

Sample

$$H_{2(d)} + \frac{1}{2} O_{2(d)} \longrightarrow H_2O_{(d)}$$
 $\Delta H = -285.85 \text{ kJ/mol}$

They tell about the exothermic and endothermic nature of the reactions and quantitative scount of heat evolved or absorbed

Why is it necessary to mention the physical states of reactant and products in a

Substances have different energies in different physical states. Therefore, heat changes all be different in different reaction, when some substance is involved in different physical

e.g. heat of formation of liquid water and heat of formation of water in vapour phases ædiferent i.e.

$$H_{z_{(0)}} + \frac{1}{2} C_{z_{(0)}} \rightarrow H_{z} C_{(0)} \qquad \Delta H = -241 \text{ fel/mor}$$

Hau + 1/2 Osla → H₂O_{th} ∆ H= ~ 285.85 kJ/mot

force, it is hecessary to mention physical states of substances in thermochemical equations.

(II)(a)Deline and explain Hour's law of constant heat summation. Explain it with examples and give its application

loked on Page 405

I lies I helps us to calculate the heats of those reactions, which accept be sormally carried out in a laboratory. Explain it lotted on Page 405

(a) What is lattice energy? How does Born-Haber cycle help to calculate the lattice energy of NaCi7 lated on Page 407

for NaCi, a closed Born-Haber cycle can be drawn as shown in figure. Thus, according

AHP = AHE + AH

Na_{cot} + 1/2 Cl_{mic} - Na'_{cot} + Cl_{tel}

the 4H' = Enthalpy of formation

AH Ne CT AH

4HL = Lattice energy

^aH_{*} = Enthalpy of conversion of Na and Cl into Na* and Cl* ions.

the this equation shows that heat of formation (ΔH_i^a), is the sum of all other enthalpies,

```
NUMERICAL PROBLEMS (Exercise)
```

Q13. 50 cm² of 1.0 M HCl is mixed with 50 cm² of 1.0 M NaOH in a glass cates The temperature of the resultant mixture increases from 21.0°C to 27.7 Assume that colorineter losses of heat are negligible. Calculate the calculate the calculate the calculate the colorineter for the reactions. The density of solution to be considered. cm 3 and specific heat is 4.18 J g^4 K^4 .

```
Volume of acid/ or base used: = V = 50 cm<sup>3</sup>
             initial temperature = T_1 = 21 \, {}^{\text{D}}\text{C}

Pinal temperature = T_0 = 27.5 \, {}^{\text{D}}\text{C}
```

Rise in temperature $= \Delta T = 27.5 - 21 = 6.5$ °C

Specific heat of water $= s = 4.18 \text{ Jg}^{+}\text{K}^{-1}$

Since density of solution (water) $= d = 1 g \text{ cm}^{-4}$

Total volume of solution = $V = 50 \text{ cm}^3 + 50 \text{ cm}^3 = 100 \text{ cm}^3$

So, mass of solution $= m = d \times V = 1 \times 100 = 100 g$.

Thus, heat evolved is given by

q = m x s x &T $= 100 \times 4.18 \times 6.5 = 2717 J = 2.717 kJ$

or q = -2 717 kJ (exothermic reaction)

Since Molanity of acid/ or base used = M = 1.0 M, therefore 1000 cm³ of solution contain HCl = 1.0 moles

 $80~\text{cm}^4$ of solution contain HGI = $\frac{1.0}{1000} \times 50 = 0.05$ moles

0.05 males of HCI react with 0.05 males of NeOH

Thus heat of neutralization is given by 0.05 moles of HCl produce heat = -2.717 kJ

0.05 moles of HCI produce heat = $-\frac{2.717}{0.05}$ = 54.34 kJ/mol

Hence heat of neutralization = $\Delta H_n^0 = 54.34 \text{ kJ/mol}$

Q14. High ratios (N_2H_4 , is a rocket fuel. It haves in O_3 to give N_6 and H_2O

The reportation is $N_2N_4 + N_2 \rightarrow N_2 + 2N_2O$. It names to D_2 to give N_2 are $N_2N_4 + O_2 \rightarrow N_2 + 2N_2O$. I. $D_2 = 0$ North is burned in a bomb coloringter. An increase of temperature 3.5% recorded. The heat capacity of calorimeter is 5.5 kJ K $^{-1}$ Calculate the quantity colored. Also calculate the D_2 calcul evalued. Also calculate the heat of combustion of 1 male of Neil.

Mass of hydrazine = m = 1 g

Heat capacity = c = 5.5 kJ K⁻¹ Rise in temperature = AT = 3.51 °C

Heat evolved during combustion is given by

 $q = c \times \Delta T$

 $= 5.5 \times 3.51 = 19.305 \text{ kJ}$

Since reaction is exothermic, therefore

q = 19.305 kJ

Molecular mass of hydrazine $(N_a H_a) = 32 \text{ g mp}^{-1} = 1 \text{ mol}$

1 g of hydrazine produce heat = = 19 305 kg

32 g of hydrazine produce heat = $-19.305 \times 32 = -617.76$ kJ/mol

Hence heat of combustion of hydrazine = $\Delta H_c^0 = 617.76$ M/mol.

 \overline{RS} . Destine (C_0H_{10}) is a motor fuel 1.80 g of a sample of octate is burned in a bomb calorimeter having heat capacity 11.66 kJ K^* . The temperature of the calorimeter increases from 21.36°C to 28.78°C. Calculate the heat of combustion for 1 g of value. Also, calculate the heat for I male of actume.

Mass of potang = m = 1.80 gHeat capacity = $c = 11.66 kJ K^{-3}$

Initial temperature = T₁ = 21.36 °C

Final temperature = T₂ = 28.78 °C Rise in temperature = $\Delta T = 28.78 - 21.36 = 7.42$ °C.

Hist evolved during combustion is given by

Q= CHAT

 $= 11.66 \times 7.42 = 86.52 \text{ kJ}$

Since reaction is exothermic, therefore

 $9 = 86.52 \, kJ$

Molecular mass of octans = 114 g mol + = 1 mole

Stace

1.80 g of octane produce heat | = - 86.52 kJ

 $86.52 \times 1 = -48.067 \text{ kJ}$ 1 g of octane produce heat= -

1.8

114-g of octune produce heat= $-\frac{86.52}{10} \times 114 = -5479.6 \text{ ke/mod}$

Hence heat of combustion of octans # $\Delta H_c^0 = -5479.6$ Li/mol

410

Q16. By applying Ham's his calculate the craftelpy charge for the formatic

o of NH_CI from NH, gas and HCI.

Sommon

Shall the will are

Office Characteristic Parts

(iii) If the heats of combination of C₂H₂, H₂ and C₂H₄ are -337.2. -68.3 and -372.8 isostories respectively, then calculate the heat of the following rescales: $C_2H_2 + 2H_2 \longrightarrow C_2H_4$

415

Heat of combustion of
$$C_2H_u$$
 is $C_2H_2 + 5/2O_{14d} \rightarrow 2CO_{24d} + H_2O_{2d} = 4H=-337.2$ keel mod [1] Heat of combustion of H_2 is $H_{24d} + 7/2 O_{24d} \rightarrow H_2O_{2d} = 4H=-68.3$ kcal raph [4] [2]

Washington and the

Graphite and diamond are two forms of carbon. The anthalog of cumbustion of symphite at 25°C is 393.51 is limit and that of diamond is 395.4 kelmod

College Charactery Part 4

420

What to meaning of the term enthalpy of ionisation? If the heat of neutralization of HCi and NoOH is -57.5 kilmol and heat of neutralization of CH₂COOH with NoOH is -55.2 kilmol. Colculate the anthalpy of ionisation of CH₂COOH.

Enthalps of ionization

thating of fontention. It is the enthalpy change when 1 mole of substance lonkes in solution to form ions under standard conditions. Example

 $CH_aCOOH_{pag} \longrightarrow CH_aCOO_{pag} + H^*_{pag}$ $\Delta H = +2.1 \text{ kJ mot}^{-1}$

Solution of Numerical Parts

Given

Heat of neutralization of acetic acid and NaOH is CH₈COOH_[eq]+Na*_(eq)+OH*_[eq] CH₈COO*_[eq]+Na*_[eq]+ H₂O

Heat of neutralization of NaOH and HO is Na $^{\circ}$ _{legt} + OH $^{\circ}$ _{legt} + H° _{legt} + OH $^{\circ}$ _{legt} + H° _{legt} +

 $H^*_{(eq)} + OH^*_{(eq)} \longrightarrow H_aO_{(eq)} \quad \Delta H = -57.3 \text{ kJ mol}^{-1}$ (2)

Regulred

Heat of igneration of acetic acid is given by $CH_{B}COOH_{(eq)} \longrightarrow CH_{B}COO^{-}_{(eq)} + H^{+}_{(eq)}$

ΔH = ?

Revening eq(2).

Writing eq. (1) as such, and then adding all these equations, we get

 $\begin{array}{ccccc} CH_{0}COOH_{(eq)} + OH^{-}\underset{(eq)}{\longmapsto} & CH_{0}COO^{-}_{(eq)} + H_{0}O & \Delta \ H = .55.2 \ kJ \\ H_{0}O_{(eq)} & & \Delta \ H = +57.3 \ kJ \end{array}$ ΔH=+573 κJ

CH₈COOH_{free} --- CH₈COO -_{red} + H*_{cep} &H = +2.1 M mol - 3

Q21(a) Define (i) Lattice Energy (ii) Atomization Snergy
(b) Draw a complete, fully tabelled Sorn Haber code for the determination of KBr.
(c) Using the information given in the table below, calculate the lattice energy of KBr.

ΔH_e = -392 κJ mol⁻¹ Christian Paris

421 ۵ السح>

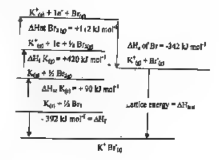
 $K^*_{ij} + Br_{ij} \longrightarrow KBr_{ij}$ secreting to Born-Haber cylice $\Delta H_{last} + \Delta H_{st}(K) + \Delta H_{t}(K) + \Delta H_{st}(B_{t_0}) + \Delta H_{st}(B_{t_0}) = \Delta H_{t}$

or $\Delta H_{int} = \Delta H_{i}$ = $\Delta H_{at}(K) - \Delta H_{at}(K) - \Delta H_{at}(B_{Y_2}) - \Delta H_{at}(B_{Y_3})$

≈ -392 - 90 ≈ 420 - 112 (-342)

= -672 kJ mol -1

日本学院の対象を





(d) free energy

(d) Neutralization

HELLO! Mr. Question here!

Multiple Choice Questions from PAST

- In endothermic reaction AH is taken as: (Laton board, 2014)
 - (c) Zero
- (b) Negative (d) May be any value la. Positive
- 2. Whenever a reaction is exothermic, then it means that: (Sujramale board, 2016)
 - (a, the heat is transferred from surroundings to the system
 - (b) the heat content of the reactant is greater than products
 - (c) the heat content of the reactants is less than those of products
 - (d) the heat is transferred from system to the surroundings
- 3. A state function which describes together the internal energy and the product of pressure and volume is called (Gayraman's bound, 2011)
 - (a) enthalips (b) internal energy (c) work Which of the following is not a state function; (makes board, 2011)
 - (a) Pressure (b) Volume (c) Temperature (d) Heat
- The Crucible which is used in Bomb Calorimeter is made up of: (Multim board, 2011)
- (a) Glass (b) Copper (c) Gold
- For the reaction, H* + OH, the change in enthalpy is called heat of (Boundplant bound.)
- (a) reaction (b) combustion (c) solution
- Standard enthalpies are measured at: (Lehore Board, 2009)
 - (a) 273k (b) 298k (c) 373k (d) all of these
- 8. For the reaction NaOH + HCI \rightarrow NaCl+H $_{\rm g}$ O, the change in enthalpy is called: Gogodia Sond, 2009) (P.G. Kleen Soud, 2011)
 - (a) Heat of neutralization (b) Heat of combustion
- (c) Heat of formation (d) Heat of reaction
- 9. The total heat content of a system is called (Labora Board, 2010)
- (a) Entropy (b) Enthalpy (c) temperature (d) internal energy
- In a bomb calorimeter, the reactions are carried out at constant. (Surgodin Board, 2020) (a) pressure (b) temperature (c) volume (d) none of these
- 11 Formation of NH_a is an exothermic reaction. However, in Haber's process temperature
- used is. @chample Board, 2020) (a) 200°C (b) 300°C (c) 400°C
- (d) 500°C (Resemblant Secret, 1010)

- (a) heat of formation of water
- (c) heat of neutralization
- (b) heat of fermation of NaCl (d) heat of reaction
- (3) Which one of the following process is an exothermic? (0.0. (then Sound 2016) (a) sublimation (b) respiration (c) Fusion
- (4. The excitheunic process is (Labora Board, 2021)
- (a) Evaporation (b) sublimation (c) respiration
- 15. Slandard enthalps change is measured at (a) 298 K (b) 273°C
- (d) bolling
- (d) 373 G 16. The enthalpy change when one mole of a substance is completely burnt in excess of oxygen is called: (Lahore Board, 2007)
 - (a) Enthalpy of atomization (b) Enthalpy of neutralization

(c) 273 k

- (c) Enthalpy of combustion (d) Enthalpy of formation
- 17. The value of ΔH being very small, the term $\Delta ({\rm PV})$ can be neglected, for the process involving (Paladobed Speed, 2010)
 - (a) Liquid and gases
 - (b) liquid and solids
 - (c) Solids and gases
- (d) None of these
- The enghalples of all elements in their standard states are: Plakes board, 2016)
 - (a) unity
 - (b) zzm
- (c) always positive (d) always negative
- 19. For the reaction NaOH + HCI \longrightarrow NaCI +H $_2$ O the change in enthalpy is called 49.6. Khan Beard, 2012)
 - (a) Heat of reaction
- **(b)** heat of formation
- (c) heat of neutralization
- (d) heat of combustion.

ismers	to Ma	dtiple (Choice	Quest.	ions fre	om Pus	a Papa	TS.
Ans	Q#	Ang	Q#	Ana	Q#	Ana	Q#	Ans
(a)	2	(b)	- 8	(a)	4	(d)	5	(d)
(선)	7	(b)	8	(a)	9	(b)	10	Jc,
(c)	12	(c)	13	(b)	14	(2,	15	(a,
(c)	17	(b)	18	(b)	19	(c)		
	Ans	Ans Q# (a) 2 (d) 7 (c) 12	Ans Q# Ans (a) 2 (b) (d) 7 (b) (c) 12 (c)	Ans Q# Ans Q# (a) 2 (b) 3 (d) 7 (b) 8 (c) 12 (c) 13	Ans Q# Ans Q# Ans (a) 2 (b) 3 (a) (d) 7 (b) 8 (a) (c) 12 (c) 13 (b)	Ans Q# Ang Q# Ane Q# (a) 2 (b) 3 (a) 4 (d) 7 (b) 8 (a) 9 (c) 12 (c) 13 (b) 14	Ans Q# Ans Q# Ans Q# Ans (a) 2 (b) 3 (a) 4 (d) (d) 7 (b) 8 (a) 9 (b) (c) 12 (c) 13 (b) 14 .c)	(a) 2 (b) 3 (a) 4 (d) 5 (d) 7 (b) 8 (a) 9 (b) 10 (c) 12 (c) 13 (b) 14 (c) 15

Detailed Explanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK-I





Their occupation

SHORT & LONG QUESTIONS FROM PAST PAPERS

EXOTHERMIC AND ENDOTHERMIC REACTIONS

Short Owestfore

- What are exothermic and endothermic reactions. Give one example of each, 10.0, 10.0, 10.0.
 Board, 2007: Labora Board, 2009: Saharaipur Baard, 2010)
- (2) Why in exothermic reactions, heat is released from the system? (Gujrenivele Board, 2018)
- Differentiate between for Compare, exciterraic and endatherraic reactions. This Search, 2007: Labore Board, 2017; D.G. Rhan Board, 2018; Asad Kashnir Board, 2012; Surgedia 2007 2012; 2018.
- (4) What is the couse for the chemical change? (Familiant Board, 2021)

SPONTANEOUS AND NON-SPONTANEOUS PROCESS

- What is a spantaneous process? Give examples. (Foliational Board, 2008; Multan Board, 2014, 2014. Gyrenucla Board, 2014.
 Give two examples of non-spontaneous reaction? (Reinstantial Board, 2009)
- (3) What are spontaneous and non-spontaneous processes? (Galranualo Board, 2010)
- (4) Spontaneous reactions always proceed in the forward direction. Give reason. OR Spontaneous are unidirectional. Give reason. (Guyrenselo Board, 2012)
- (5) Differentiate between spontaneous and non-spontaneous reactions. (Palastebod Board, 2011: Baselphad Board, 2011: D.O. Rhom Board, 2012: Multan Board, 2012: Outrampole Board, 2012: Gargodha Board, 2014)
- (6) Spontaneous reactions are exothermic in nature. Explain. (Gujrameda Board, 2011) Lowe Quantions
- Differentiate between (!) Spontaneous and non-spontaneous reaction (!) Exothermit and Endothermic reaction (Submedger Board, 2009)
- (2) Differentiate between (1) Internal energy and enthalpy (III) Exothermic and endothermic reaction. (Ravelpinel Board, 2009)

STATE, STATE FUNCTION, SYSTEM, SURROUNDING, BOUNDARY, INTERNAL ENERGY Short Countiens

- (1) Define state and state functions with examples. (Lehen Board, 2008; Sufranuela Board, 2000)
- (2) What is meant by state function, Explain with top examples. (Sergodia Board, 2008: Sujnonwald Source, 2008: Submaripur Board, 2012: Multiur Board, 4012: Labore Board, 2010, 2013, 2014)
- Bonnardyn Board, 2012: Multon Board, 2012: Labore Board, 2010, 2013, 2014

 (3) Define for describe or differentiate) system and surrounding with one example.

 (Outraneous Board, 2009: D.G. Khan Board, 2009: 2011: Roll2: Falactobed Board, 2010: Salesmines

 (4) Define system and state function. (Sargadha Board, 2012: Surpodha Board, 2018)

 15) Tennarothura is a state function.
- (5) Temperature is a state function but heat is not. Why? (0.0. Khen Found, 2008)
- (6) What is state function? Prove that Internal energy is a state function. (7) (Releasiper Beard, 2010)
- (8) Define Internal Energy, (Malten Board, 2012)
- (9) What do you know about internal energy of a system? Give excursols (Saharelyer Booth 2012: Makes Soort, 2012: Laiera Boord, 2016)

 Define the larm Vacule' and convert 15 calories to joule (Sargedha Boord, 2015)

College Characters Parts

123

MET LAW OF THERMODYNAMICS ENTHALPY

Short Guretlana

- Shot West-same
 1) Define first law of thermodynamics. How it is represented mathematically? (Lake
 2017: Feledakad Goard, 2009: Multan Beard, 2016: Resembled Board, 2011)
- What is the significance of the equation $\Delta F = q + \omega T$ (Remarkable Be-
- (3) Explain the term enthalpy (Later Seed, 2018)
- (6) Prove that change in enthalpy is equal to heat of reaction. (8.6, 10cm Sect., 2010) OR Prove that $q_0 = \Delta H (D.G. Khow Board, 2012)$
- State why, $\Delta H \approx \Delta E$ in case of liquids and solids, μ_{m} nipindi Beard, 2007)
- (6) Differentiate between internot energy and enthalpy (Fancished Board, 2012; in 1964, 2012)

- (1) State and explain first law of thermodynamics. (Labore Board, 2009)
- (2) What is first law of thermodynamics. How does it explain that $AH=q_{\mu}$ (i.e. Rims &c.
- (3) Discuss first law of thermodynamics and prove that $\Delta E = q_s$ (0.6. Non-Board 2019) OB State first law of thermodynamics Prove that $\Delta E = q_s$ (Labore Board, 2013)
- (4). Explain enthalpy and prove that $\Delta H=q_0$ (Laters Secré, 2010; Mailon Board, 2013)
- (5) Define system and surrounding. Also prove $dH=q_{\rho}$ (Federal Bosed, 2010)
 - (6) Prove that $\Delta k_i^2=q_p$ (Labora Bourd, 2007; Outroweds Bourd, 2020)
 - Of Prove that change in Internal energy of a system at constant volume is equal to heat obsorbed by the system. (b.6. Khan board, 2007) OR Prove that $\Delta E = q_{\phi}$ (Labora Board.
- (8) Prove that: $q_p \Delta E = q_q$ (6) $\Delta H = q_p$ (Gajvanuela Board, 1903: D.G. Khan Board, 2011)

ENTHALPHY OF REACTION, ATOMIZATION, PORMATION, MELITRALIZATION, COMBUSTION, SOLUTION

1 Define ALC

ofine AH? Can It be negative. Why? (Matun Sound 2008)

- What is mount by standard enthalpy of atomisation? Give example (Pstechbut Bound, 2006.

 And Bushirt Board, 2012 Resolved Board, 2013: Labore Board, 2013: Surgodio Board, 2014)

 What is meant by standard enthalpy of neutralization? Give the example (D.O. Klum Board, 2008, Public Board, 2008, 2018). Surgodio Board, 2009. Resolved 2012, 2013: Labore Board, 2014: Bujranenie Beerd, 2014; OR Enthology of neutralization is for an acid and have. Explain with example: (Bukewilper Bood, \$41.1)
- (4) Comment that enthalpy of neutralization is merely the heat of formation of one mole of liquid water it above Seard, 2005 OR Enthalpy of neutralization for any strong acid with (5) Dignet Explain with examples
- - .(i) entirely, of compassion (Mulei Board, 2012, D.O. 10on Board, 2009, 2018, Out
 - (II, enthalpy of atomisation and enthalpy of neutralization (Seguido Board, 2016)
 - (iii) alomization energy and lattice energy? (Handpind Board, 2014)
- (to, standard enthalpy of fermation (Labore Board, 2012)





\$24

(v) mandard enthalpy of colidion (Osternational Seed, 2008; European Roset, 2018) (vi) enthalpy of combustion and enthalpy of solution (Revelokal Seed, 2018) (vii) enthalpy of formation and enthalpy of atomisation. (Februaries Seers, 2007)

(1) Define the following. (New-Little 1-4, 2007)

(i) Standard enthalphy of neutralization with example. (N) Standard entholphy of combustion with example.

(2) Define the jollowing:

(i) System /Cogine le Board, 2011)

(il Surroundings Patrolabed Souré, 2007: Oujran da Board, 2011)

(ii) Standard enthalpy of atomization (Principle Search, 2007; Outrasselle Search, 2011)

(iii) Standard enthalpy of solution. Parking Sours, 2007: Gujranuscie Bourd, 2011)

MEASUREMENT OF ENTHALPY OF REACTION

Bert Constinu

(1) For what purpose bomb calorimeter is used? (Regodie Seent, 2007)

Less Counties.

(1) What is enthalpy of a reaction? Name two methods for the determination and amids. any one method. (Surjoins Souré, 2009)

(2) Define enthalpy of reaction. How the enthalpy of reaction is measured by size colorimeter? (Surgette Board, 2012) Lahon Board, 2012)

[3] Define enthalpy of neutralization. Also discuse gloss calc "meter in cletail? (Guy-

Saura, 8818) on State the construction and worlding of glass colorimeter (Gramm

(4) Describe how the enthalpy of combustion (dHe) of a substance is measured by bomb colorbrater? (Fatebaad Surd, 2015, 2012; Receiphell Board, 2014)

HERE'S LAW OF CONSTANT HEAT SUBDIATION, BORN-HABER CYCLE

(1), Sinte the Hears law of coretant heat nonmation (Labore Board, 2015; Falcatobal Bo (2) Justify that heat of formation of compound it turn of all the other enthalpies. (Liberation of compound it turn of all the other enthalpies. (Liberation)

(3) What is Born-Haber cycle? What is the main advantage of "Born-Haber cycle" (0.6. Illustrated 2007 Males Bound, 2013)

(4) Draw o fully labelled "Barn-Haber cycle" for the formation of NoCl (Males Bound, 2009)

(1) Destrict or Define or State "Heat's law of constant heat numeration" with an example frequency flam, 1906; 36-bit Band, 3918, 3919 Sargadha Band, 2007, 2018, 2018, 3918 (2) Define Lattice Energy, and Born-Haber cycle Host lattice energy is measured by 80°F Hober cycle Write assessing of Assessing Page Host lattice energy is measured by 80°F Hober cycle.

Hober cycle. Write equation of different anthology changes in the formation of NaCl form its electronia, Medicinalism Samed, 2011 Surpriller Board, 2013)

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Den 50 at s. 8 a

A ST AND ASSESSED AS (a) PV (b) PAV (c) APV (d) Appending out of constant opening of the property o

in a company comment of the constant pressure (d) constant temperature (d) a, b, a condition which of the following is the correct equation?

Which of the lobowing is the correct equation? If $\Delta E = aw + \Delta q$ (d) note Europoration of water in [b] $\Delta w = \Delta E + \Delta q$ (c) $\Delta E = aw + \Delta q$ (d) note Europoration of water in [b] on exothermic change (b) a chemical reaction (c) an endothermic change (b) a process where no heat changes occur Hassiaus status, energy, of formation of a compound (e) depands on path (b) does not depand on path (c) both e end b are accord (d) note of the above Hassia data with

Hear a lew clean with

a) changes in heat of reaction
(c) equilibrium constant
(d) influence of pre-

(d) influence of pressure on volume of a gas

Combustion of methans

| Section | Communication | Co

remure is equal to (d) H

 (a) a bootstand by a system of constant print (a) ΔΕ (b) ΔΕ (c) Ε (c)
 (b) ΔΕ (c) ε (c) ε (c)
 (c) ε (c)
 (d) ΔΕ (d) ΔΕ (d) ε (d)
 (e) ε (d)
 (f) ε (d)
 (g) δΕ ne whether it isless place in one step or in several steps. This (a) first law of thermodynamics (b) law of thermochemistry (c) Heav's Levy (d) Record's Law

 ΔH_{ζ}^{0} is the standard enthalpy when

[8] 3 mote of a compound formed from its stammerts (b) 1 mote of an increased formed fines its ions.

[9] 1 mote of a compound formed from its stammerts (b) 1 mote of an increased formed fines its ions.

[9] 1 mote of a compound burnt in recome of oxygen complaintly.

[9] 1 mote of a compound burnt in recome of oxygen complaintly.

[9] 1 mote of a compound burnt in recome of oxygen complaintly.

[9] 1 mote of a compound burnt in recome of oxygen complaintly.

[9] 2 mote of a compound burnt in recome of oxygen complaintly.

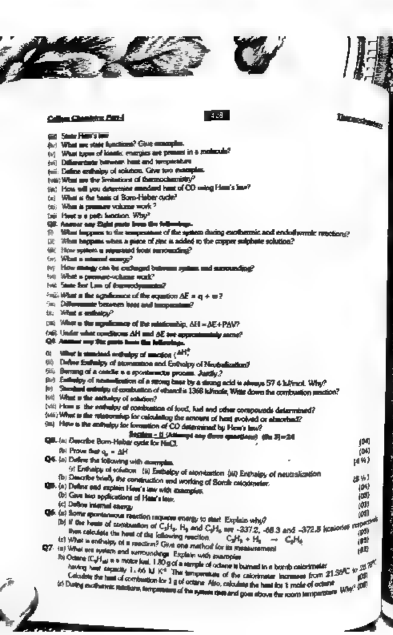
[9] 2 mote of a compound burnt in recome of oxygen complaintly.

[9] 2 mote of a compound burnt in recome of oxygen complaintly.

[9] 3 mote of a compound burnt in (b) 1 mote over the compound of the compo (ii) Instabilities of creation (b) rotational rection (c) vibrational motion (c) Standard enthalty of combanion of H₂in 200.8 kJ mod ² (b) which is the standard enthalty of loyer (a) + 200.8 kJ mod ² (c) Zero (d) -21 kJ mod ³ (d) -200.8 kJ mod ³ (d) Zero (d) -21 kJ mod ⁴ (d) -21 kJ mod (4) - 218 년 (6)

On of Questions 2,3 and 4, Militar any TWENTY TWO(22) short assess. While unline assessment with a property of the control of

May a list necessary to memory physical pages of education is a theoretic Discoverable between approximated and new-special recommendate process.



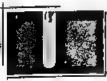
hapter 8 CHEMICAL EQUILIBRIUM



Electronic pH meter showing pH = 6.82







BiOCI + 2HCI



CONTENTS

Chapter-8

Chemical Equilibrium

REVERSIBLE AND IRREVERSIBLE REACTIONS

State of chemical equilibrium Law of mass action Equilibrium constant expressions for some resettons Relationsh ps between equilibrium constants

Applications of equilibrium constant ce-chatelier's principle

APPLICATIONS OF LE-CHATELIER'S PRINCIPLE IN INDUSTRY

Synthesis of ammonia by Haber's process Synthesis of 30:

IONIC PRODUCT OF PURE WATER IONIZATION CONSTANTS OF ACIDS (Ka) IONIZATION CONSTANT OF BASES (Kb) COMMON ION EFFECT

BUFFER SOLUTIONS

Buffer capacity EQUILIBRIA OF SLIGHTLY (SPARINGLY) SOLUBLE IONIC COMPOUNDS (SOLUBILITY PRODUCT)

Applications of solubility product

Objective and short enswer, questions (exercise) Numerical problems (exercise)
Past Papers MCOs and Short Ouestions
Tast your skills

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TURSBLE AND IRREVERSIBLE REACTIONS

lardre Q4. (a) Explain the terms "reversible reaction"

arzable Rem tirut

The reactions, soleich can proceed both in forward or well as in backward direction, are salled souls reactions. anole

The reaction of N_2 and H_2 to form NH_3 is a reversible reaction. This reaction occurs at bit in the presence of Iron as catalyst and under high P

At start, N_x and H_z react to form NHs. However, after sometime NHs also decomposes to Nz and Hz because conditions are favourable for both forward as well as for a Thus, it is a reversible reaction.

The meantion mixture contain, all the three substances i.e. No. He and NHs.

terribie la action totilons, which take place only in one direction, are called irrepresible recessors. Clambes which take place only in one arranson.

Clambes reactions can have place in both directions. However, for some cases, reverse lefter but. don is very small

0 Pacifor of Na with H_aO produce NaOH and H_a

2Na + 2H₂O --- 2NaOH + H₃ A room, temperature, reverse reaction is negligible. It is an impressible reaction.



e-Charles - Paint

- when the case of H_{α} and O_{α} to form $H_{\alpha}O$ in the presence of electric speak 2H₂ + O₂ ---- 2H₂O
- At room temperature, this seaction is negligible in reverse direction. If H₂ and 0 are mixed as correct proportion than no reactants are left behind.
- However, at \$500°C, H_eO decomposes back to H_e and O₂. Thus revene stark

Hence, at low temperature, this reaction is not reversible. It is called impumble warten.

radicatact between present	te und irreperatble Rea	ctions
Recessible reaction	Irreversit	de renetem
The reactions, which can proceed both in licrosist as well as in backward direction, are called reversible seachons.	The reactions, which one directions are careactions.	
to these reactions, a dynamic equilibrium is established Catherally, both secutarits and products are greenest at equilibrium state.	2 bi these reactions, e established. 3 In these reactions, o are almost consume	ne or more space
Extensiones Ny _{top} + 3H _{Red} et = 278H _{Red} 25O _{Red} + O _{Red} et 25O _{Red}	Examples 2Na + 2H ₂ O → 2	NaOH + H

Exercise Q4 to Explor the come "state of equilibrium"

MARCH CHEMICAL LOCHEROM

The state of councilide contains in which rate of for of contain reaction to called characted aguithment.

Software process reaction

or the conforming A & & march to give produces & this is the early sentilled tolera

e as soon as some products are formed, the per reaction starts at slower rate

As time passes, forward reaction slows down and ess reaction speeds up and after some period rates of good and reverse reaction becomes equal and thus a partiet equilibrium is established. At this stage centrations of reactants and products become

thas been shown in the graph. The concentration

got running parallels to the time axis represents the equilibrium state

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H_{tel} and I_{tel} seacts at 425°C to form HI_G.

equilibrium, the concentration of $H_{\rm e}$ and $I_{\rm e}$ and H becomes constant. These are not enged with time and a dynamic equilibrium is established.

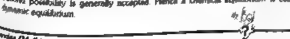
When rising curve of products and laking curve of transits becomes parallel to time axis, equilibrium is arbithed as shown in the fig.

Further, same concentration is obtained whether the then is carried out by reacting H_{deg} and I_{per} or by transporting Hie

I thous that there may be two possibilities

- Either all the reactions have stopped in the when and the system becomes stationary
- * Or both forward and reverse reactions are occurring at the same rate

Second possibility is generally accepted. Hence a chemical equilibrium is called an a



redex Q4 (ht): And explain the law of mean action and drive the expression for the equilibrium MAN K

SEIN MAYS AL SHIN

A Couldbarry & P Strange green in 13664 in conse









The rate of which a substance reacts is directly proportional to it matter many rate of a reaction is directly proportional to the product of active masses of

Active mass means concentration of mactants and products in mos den in a dissolution.

Destination of K. Expression

Consider a general reaction

The active masses of A , B ,C & D in terms of mole don 3 are represented by [A], [B][C] & (D) respectively.

According to law of mass action

Rate of forward reaction $(R_t) = \alpha (A) [B]$

and Rate of backward reaction(R_c) α [C [D]

or
$$R_i = k_i[C][D]$$

Where $k_f = rate constant for forward reaction$

k, = rate constant for backward reaction

At equilibrium statu

$$R_i = R_i$$

$$k_{i}[A][B] = k_{i}[C][D]$$

$$\frac{k_{i}}{k_{i}} = \frac{[C][D]}{[A][B]}$$

$$\frac{K_1}{K_2} = \frac{(\Delta)(B)}{(\Delta)(B)}$$

$$R_c = \frac{[C_n]D]}{[A][B]}$$

Where K_e= Equilibrium constant

K is the ratio of two rate constants.

The subscript 'c indicates that concentration are expressed in terms of

Conventionally, concentrations of products are written above and concentrations of Mactanta below

$$R_c = \frac{[Products]}{[Reactants]}$$
 or $R_c = \frac{Rate constants for forward step}{[Pate constants for backward step]}$

At a given temperature, value of K, is independent of the initial concentration.

(ii) The value of K, varies with hemperature

College Characteristics Ports 1

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Rora more general reaction

The co-efficients appears as exponents of the concentration terms in the equilibrium populant expression.

$$..e., K_4 = \frac{[C]^c[D]^d}{[A]^c[B]^b}$$

A state of dynamic equilibrium helps to determine the composition of the reactants and products at the equilibrium,

(mrs of Lipsidderson Constant

Julis of Ho depend upon the number of moles of reactants and products involved in the пофрант

J. Resetton setthout change in number of moles

For such reactions $K_{\mathbb{C}}$ has no unit

$$K_0 = \frac{|CH_3COOC_2H_3]||H_1O|}{|CH_3COOH||C_2H_5OH|} = No \; Units$$

2. Repations with change in number of moles

Por such reactions units of K_C varies from reaction to reaction

$$K_{C} \triangleq \frac{N[H_{3}]^{2}}{[N_{2}][H_{2}]^{3}} = \frac{\left[\text{moles dm } 3\right]^{6}}{\left[\text{moles dm } 3\right]^{3} \left[\text{moles dm } 3\right]^{3}} = \text{mol}^{-3} \, \text{dm}^{-6}$$

Example I The following reaction was allowed to reach the state of equilibrium.

2A + B - C

The initial amounts of the reactions present in one dm' of solution were 0.50 mole of A and 0.60 mole of B. At equilibrium the amounts were 0.20 mole of A and 0.45 mole of B and 0.15 male of C. Calculate the equilibrium constant Kg



on Chemistry: Parts

Chamiral South

Q6 (a). Write down Ke for the following reversible reactions. Suppose that the volume of reaction indure in all the cases is "V" dim" at equilibrium stage.

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EQUILIBRIUM CONSTANT EXPRESSIONS FOR SOME REACTIONS

1 Formation of I stee trave an Organic Acid and Alcohol: (Aquisaus Phase Reaction)

Alcohol and acid reacts to form exter and water in the presence of small amount of mineral acid as catalyst 100+

The progress of reaction can be noted by finding out the concentrations of acetic acidal regular interval. A small sample of reaction mixture is taken out and amount of acid a determined by titrating it with standard NeOH

Let initial cond of acid is 'a mol/dm' and that of ethanol is 'b' mol/dm' and volume of container is 'V' dm'. If at equilibrium, 'x' moles of spetic acid-reacts with 'x' moles of ethanol to give 'x moles of ethyl acetate and 'x' moles of H_1O , then

According to Law of Mass Action

$$K_{s} = \frac{\text{[CHaCOOCsHs]{HsO}}}{\text{[CHaCOOH][CsHsOH]}}$$

$$K^{z} = \frac{\begin{pmatrix} A & A \\ B & A \end{pmatrix} \begin{pmatrix} A & A \\ A & A \end{pmatrix}}{\begin{pmatrix} A & A \\ A & A \end{pmatrix} \begin{pmatrix} A & A \\ A & A \end{pmatrix}} = \frac{(a - A)(p - A)}{A_{x}}$$

The final K_c expression contains no V term. Hence volume changes for president changes does not affect K_c or equilibrium position of reaction.

Chemistry: Part-

2. These lation of PCI, (Gas Phase Reaction) PCI, PCI, + CI,

If initial come, of PCl₀ is 'a' and volume of container is $V \, dm^2$ and at equilibrium \times moles of PCl₀ and \times moles of Cl₀, then

9	PCI,	PO,	+	Cl.
Julitlas conc.	a	Q		0
(jn moles) (1=0)				
Equilibrium conc	a x	×		x
(In moles: (Lest_)				
Equilibrium conc./V	(a x)	к		x
Edimonant court 4	V	√		V

(mol/dm⁻³)

According to law of Mass Action

$$\begin{split} K_c &= \frac{\left[PCI_3\right]\left[CI_2\right]}{\left[PCI_3\right]} \\ K_c &= \frac{\left(\begin{matrix} x & \\ V \end{matrix}\right)\left(\begin{matrix} x \\ v \end{matrix}\right)}{\left(\begin{matrix} a-x \\ v \end{matrix}\right)} = \frac{x^2}{\left(\begin{matrix} a & x \end{matrix}\right)V} \end{split}$$

The final K_c expression contains V term. Hence K_c and equilibrium position of reaction in affected by volume and pressure changes.

iccomposition of $N_{ m c}$	D. (Gas I	Thosa Read	dion)
	N ₂ O ₄₉₀		ZNO
initial some	a		Q
(In males (t=0) Equilibrium conc. (In moles 't=)	z -x		2x
Equilibrium conc./V	(a <u>x)</u>		2 <u>x</u> √
(motelm -s K _e ls given by			
$K_c = \frac{ NO_2 }{ N_2O_2 }$			



College Charles Call

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$$K_{x} = \frac{4x^{2} - V}{|a-x| - V^{2}} = \frac{4x^{2}}{|a-x| \cdot V}$$

The final K-expression contains V temp. Hence K- is affected by volume and treate honces.

4. Stath, six of NH, (Gos Phose Resultion)

	Name	+ 3H _{2M}	
salfal conc	a	ь	0
(in moles) (t=0)			
Equilbrium conc.	ах	ъ 3х	2x
(in moles (fut)			
Equilibrium conc./V	$\frac{(\mathbf{a}-\mathbf{x})}{V}$	(b-3x) V	<u>2x</u> ∨
(mol / das *)			

 K_c is given by $K_c = \frac{[NH_3]^3}{[N_2][H_2]^3}$

$$\begin{split} & R_c = \frac{\left(\frac{2x}{V}\right)^2}{\left(\frac{b-3x}{V}\right)\left(\frac{b-3x}{V}\right)^3} \\ & R_c = \frac{4x^2}{(a-x)V^2} \times \frac{V^4}{(b-3x)^6} = \frac{4x^2V^8}{(a-x)(b-3x)^6} \end{split}$$

Thus, co-efficient of balanced chemical equation determines whether the volume term will appear in numerotor or denominator

5. Description of III (Gas Phase Receitors)

	21-11	-	He +	l.
Initial Conc (moles) (t=0)			0	ō
Equilibrium Conc. (moist) (1=t_j)	(= -2x)		×	ĸ
Equitorium Cons.	$\frac{(a-2x)}{V}$		ž	×
(metida di	V		Ÿ	ŷ

refine Chambire Per-l

According to Law of Mass Action

$$K_{c} = \frac{\left(\frac{\ln x}{|E|}\right)\left(\frac{1}{|E|}\right)^{2}}{\left(\frac{x}{V}\right)\left(\frac{x}{V}\right)}$$

$$K_{c} = \frac{\left(\frac{x}{V}\right)\left(\frac{x}{V}\right)}{\left(\frac{\alpha}{2x}\right)^{2}} = \frac{x^{2}}{\left(\alpha - \frac{x^{2}}{2x}\right)^{2}}$$

The final K commession contains no V term. Hence volume changes or these changes does not affect K or equilibrium position of reaction.

6. Formation of HI (Gos Phose Beaction)

			- Dance	O P	
	H ₂	+	J _n		24
heitiel Conc. (moles) (1→5)	*		ь		0
Equilibrium Conc. (males) (f.e.()	(a-x)		(b-x)		2x
Equilibrium Conc	(0 - X)		(b x) V		$\frac{2x}{V}$

(molétim -*)

According to Law of Mass Action

$$K_{t} = \frac{(H_{2})[t_{2}]}{(V)^{2}}$$

$$K_{t} = \frac{\left(\frac{2x}{V}\right)^{2}}{\left(\frac{\sigma - x}{V}\right)\left(\frac{b - x}{V}\right)^{2}} = \frac{4x^{2}}{(a - x)(b - x)}$$

The final K- expression contains no v term. Hence volume changes for manufactuation, standard does not affect K- or equilibrium position of meetion.

Exercise Q7. (a) Write down the relationship of different types of equilibrium constants. i.e. $K_{\rm F}$ and $K_{\rm F}$ for the following general reactions.

fichtionships Butteren Equilibrium Constons

For a general reaction



Ke for this seaction can be written as

$$K^c = \frac{[C]^c D[^d]}{[A]^d [B]^d}$$

or Simply
$$K_c = \frac{Cc^c \times Cp^d}{Ca^0 \times Cp^0}$$

where $C_{c} = C_{b} = C_{a}$ and C_{b} are the molar concentrations of C, D, A and B respectively, If the reactants and products are ideal gases then molar conc. of each gas is proportional to its partial pressure. When the conc. are expressed in terms of partial pressures, the K x alven as

$$R_p = \frac{P_0^c \, P_0^d}{P_A^d \, P_0^d}$$

Where $P_a^* = P_b^* = P_b^*$ and P_b^* are the partial pressures of C, D, A and B at equilibrium

ilelatmiship Between K. And K.

It is given by

$$K_p = K_c \times (RT)^{trr}$$

Where

R = General Gas constant

T = Absolute Temperature

 $\Delta n = \text{change in no of moles in going from reactants to products.}$

If the number of moles of reactants and products are equal for a gaseous reaction then values of K_C and K_F are same, since $\Delta n = 0$.

Example 2

N, and H, e ins to give NH₂. Calculate K_{p} for the synthesis of NH₂. When $K_{C} \simeq 6^{\times}$ at 500°C

2NH_a (D.C Visan Board, 2011. Gujranunia Board, 2010, 2014: Lahere Board, 2013. Multan Board, 2013: Repairing Board, 2018 $N_0 + 3H_p \Rightarrow$

Solution

 $K_c = 6 \times 10^{-6}$ T = 500°C + 273 = 773 K R = 0.0821 atm dzn³ znal - 1K-1

Change in number of moles = $\Delta n = 2$ 4 = -2

Sunce

Colors Chamistry Pari-1

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 $K^b = K^c (RT)^{ba}$ $K_p = 6 \times 10^{-2} (0.0821 \times 773)^{-2}$

6×10⁻² $K_p = \frac{G \times 10}{(0.0821 \times 773)^2}$

 $K_p = 1.5 \times 10^{-9}$ Here Ke is less than Ke

The reactions, which occur with decrease in number of moles, have smaller Ke than K. and vice versa.

APPLICATIONS OF EQUILIBRIUM CONSTANT

Value of equilibrium constant is specific and constant at constant temperature. It can be used for following applications

- 1 Prediction of direction of reaction
- 2 Extent of chemical reaction
- 3. Effects of various factors on chemical equilibrium (Le chateller's principle)

Exercise Q9;

Explain the following two applications of equilibrium constant. Give examples.

(f) Direction of reaction

1 Prodletion of Direction of Reaction

If value of K_C is known for a particular reaction at a given temperature, then for a particular reaction at a given temperature, then

before reaction attains equilibrium) can predict the direction of reaction.

There may be three cases

Case - I f [product] < Kc [reactant]

In this case, more product is needed to regain equilibrium. Hence reaction occurs in ferverd direction

Case | [product] > Kc ,reactant

In this case more reactant is needed to regain equilibrium. Hence reaction occurs in backward direction





Coffine Classification (Cort.)

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Changing Paymented

Cove - Ili (f (product) (reactent)

in this case, reaction is at equilibrium and no more reactant or products are formed

Establication reaction between ethanol and acetic acidinae carried out by mixing definite anounts of ethanol and acetic acid dengatiff some mineral acidins a catalyst. Sample m out of the reaction mixture to check the progress of the esterification next denome in one of the samples drawn ofter time t. the concentrations of the species nere found to be [CH₂COOH]=0.025 scalidm², [C₂H₂OH]=0.032 malign², [CH₂COOC₂H₂]=0.05 scalidm² and [H₂O]=0.04 malign² Find and the direction of the reaction if R, for the reaction at 25°C to 4. (Sorgerdina Board, 2014)

The exterification reaction is obeyn as

 $\mathsf{CH_{3}COOC}_{1}\mathsf{H}_{2} + \mathsf{H}_{3}\mathsf{OH} \xrightarrow{\mathsf{E}^{*}} \mathsf{CH_{3}COOC}_{2}\mathsf{H}_{2}^{*} + \mathsf{H}_{3}\mathsf{O}$

Since volume is same for all substances, therefore, the K_c for the reaction will be

 $K_c = \frac{[c H_1 cooc_2 H_6][H_3 O_2]}{[c H_3 cook_1][c_2 H_3 O_3]}$

The [product]/[reactant] ratio will be

 $\frac{0.05 \times 0.04}{0.925 \times 0.032} = 2.5$

Since producti (reactant) < Kc (i.e. 4), therefore, the reaction will proceed to forward direction

Explain the following two applications of equilibrium constant. Give examples (fi) Extent of reaction

2. Astron of Chemical Beaution

Value of Ke can also tell the extent of reaction. e. how much reaction have occurred? There may be three cases

Case - . . Value of K. Very Large

Very large K, value shows that reaction is almost complete.

Thus at equilibrium there will be very high conc, of products and very low certified reactants

Example

20, 30,

Equilibrium constant for this reason is 10^{64} at 25° C

It shows that at noom temperature, O₃ is highly unstable and it almost completely decomposed to O₂.

Charleton Part-I

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Case - II : Value of Ke Small

Small K, value shows that the reaction does not proceed appreciable in forward reaction. Generall : Value of K. Very Small

It shows a very little forward reaction. Thus at equilibrium, there will be high patemitation of reactants and very low cone of products Example:

 $2HF_{i\phi} \longrightarrow H_{i\phi} + F_{z\phi}$

Equilibrium constant for this reaction is 10-18 at 2000°C.

it shows that at 2000°C, HIP is stable and it is only slightly decomposed to give product

3. effect of Changing in Conditions in Chesical Equilibrium

Equilibrium constant and position of equilibrium are two different things. K, is equilibrium constant. If has definite value at a civen-temperature, However producti ratio (reactant

called position of equilibrium. It can be changed by changing conditions. e.g. temperature passure and concentration etc.

like is large, the equilibrium lies to the right. If K, is small, the equilibrium sees to the left,

H-CHATELIER'S PRINCIPLE

(a specie is applied to a system at squilibrium, the system will not in such a way so as to sully, as far as possible, the effect of that stress.

A system cannot completely cancel the effect of change. It only minimizes the effect of change Le-Chateller's principle tells about the position of equilibrium and composition of Physical and chemical equilibriums.

Beet of Change in Concentions

Consider the hydrolysis of BiCla

BiCl_k + H₂O ==== BlOCl + ZHCl

Ke is given by

 $K_{c} = \frac{BOC()[HCt]^{3}}{2}$ [BICI:][H:O]







Color Charles Date

2.24

Chemical English

Solution of BiCl₆ is <u>depute</u> due to <u>formation of BiCCl</u>. Now if HCl is added to this system, the equilibrium will be disturbed in order to regain equilibrium, the reaction and solution becomes clearer.

On the other hand # H-Q is added to the system, # will shift the equilibrium in forward direction and the solution will again becomes cloudy.

Thus following are the effects of changing concentrations

- Adding reactors or removing product favours forward reaction.
- Adding product or removing reaction favours backward reaction.

But value of K_c remains constant. Hence, if one of the product is removed continuously the yield of reversible reaction is increased.

This concent of character concentration is applied in common ton effect.

Figer of Change in Pressure of Valuar

An increase in pressure on an equilibrium system will move the reaction in a direction of decrease in mounter of moles i.e. decreased volume I and vice versa.

The change in pressure can only effect those reactions which involve votume changes.

Generous reactions involve votume changes, therefore, they are greatly affected by pressure.

Example.

In this reaction, number of moles of reactants are <u>decreasing</u> from reactants to produce Γ . Thus as equilibrium volume of reaction mixture will be <u>less</u> than the volume of reaction taken. Thus, <u>increasing P</u> will shift the equilibrium in <u>forward direction</u> ...e. towards gestion when but value of K_0 remains constant.

Description (Efficient of Volume on Equilibrium Pasition).

Example: Formation of SO.

The mathematical K_c expression for

$$|g_{ab}| = \frac{4x^3 V}{(a - 2x)^3 (b - x)}$$

Where V = volume of seaction mixture at equilibrium.

 $\mathbf{n} = \text{Initial concentration of } \mathbf{SO}_{\mathbf{n}}$

b = Initial concentration of O_a

Committee Part !

4.5%

Chemical Equipment

 $x = Moles of O_3$ reacted at equilibrium

It shows that if P is decreased then V is Increased, therefore, is most decrease to maintain constant value of K_C. Hence reaction proves in <u>backward direction</u>.

Similarly if \underline{V} is decreased by increasing \underline{P} then K_0 is decreased. Thus reaction will move in forward reaction to maintain constant value of K_0 .

Other Examples

Its mathematical K_{C} expression is

$$\kappa_C = \frac{x^2}{(a-x)V}$$

Its mathematical $K_{\mbox{\scriptsize C}}$ expression is

$$K_{iC} = \frac{4x^2}{(a-x)V}$$

Both these reactions have volume in the denominator it is because no of motes of product are greater than no, of motes of reactants

Thus if P is increased then V is decreased, therefore, X, must also decrease. Hence, the reaction will proceed in backward direction to maintain constant value of K_C .

Pressure will have no effect on the reactions, in which number of moles of reactants and products are same, e.o.

1 moje 1 moje 2 mole
Similarly, reactions of <u>liquid and solids</u> are also <u>not affected by pressure</u> changes, because they do not involve volume changes

Difficet at Temperature

Considering heat as a component of system, an increase in temperature adds heat to the system, while decrease in temperature removes heat from the system. Hence according to a Chatter's principle

- Increase in temperature favours Endothermic reaction, while
- Decrease in temperature favours Exothermic reaction.

Value of K_c changes with change in temperature. It is because equilibrium shifts without addition or removal of reactants or products.



Chemical Englishing

Exercis 1 . Fermation of CO. from CO

CO₂₀ + H₂O₄₀ CO₂₀₀ + H₂₀₀ ΔH = -41.84 κJ/mol

it is an exothermic reaction, hence decrease in termograture fewours forward reaction and

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By decreasing temperature, heat of the system is lost. Thus more CO and $H_{\rm g}{\rm O}$ will react to produce more heat. Hence the reaction will go in forward reaction. If this system is bester, then the reaction will go in backyant reaction.

Exemple 2 : exhability of Salta

Colombia Patri

Consider the solubility of 10

King ____ King

 $\Delta H = +21.4 \,\text{kJ/mol}$

It is an endothermic reaction. Hence increase in T favours forward reaction.

Hence more and more sait is dissolved by increasing T. While cooling results in hackeged process. Thus Ki_{es} creatilizes out.

For some sales, heat of solution is almost gazo, For such sales change in T has no effect e.g. formation of NaCles

If exceptances have negative heat of solutions e.g. LICI , LyCO₃ , then their <u>solutions</u> decreased by increased temperature.

the specification of

A constget to σ and common which changes the rate of σ chemical reaction without tieff failing command in the process.

Thus catalyst has <u>no effect</u> on the equilibrium composition of the system. It simply stockers, the rate of forward and backward reaction. Thus, it only decreases the first if many the equilibrium state.

A complete some ing activation ensure of the reaction by giving new path to the reaction

Some storms of Led intelliers termetale in it dustry. Contenses of Animonists, Robert Process

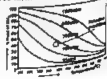
The process of synthesis of NH₄ was introduced by German chemist F. Haber in 1933.

N_{No.} + 3H_{No.} —— 2NH_{No.} K_c is given by

 $\Delta H = -92.46 \text{ kJ/med}$

 $K_C = \frac{[NH_2]^d}{[N_2][H_2]^d}$

According to Le-Chateller's principle yield of $NH_{\rm B}$ can be increased as follows.



Contract Party

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والمالية ليشطان

By continuous national of NH₂ from the reaction mixture reaction moves in forward direction.

g Since there is decrease in number of moles in going from reactants to products. Hence increase in pressure favours synthesis of NH₃. The pressure used is 200-300 atm.

Since reaction is <u>excitenmic</u>. Hence, <u>decrease in temperature</u> favours forward reaction. The temperature used in 400°C

Hence high presence. Itse temperature and continuous removal of NH₂ will give best at NH₂.

eg. Value of K_0 at 200 K is 7.17×10^{58} , and at 600 K it is 140^{-11} for the temperature on K. for attentions scallesses

Decrease in value of $K_{\rm C}$ shows that increase in T decreases a synthesis of $NH_{\rm S}$

on Conditions for the synthusts of NH₃

is actual process, following optimum conditions are used

Pressure = 200 - 300 aim

(9) Temperature = 400°C (673 fc)

(11) Catalyst = Pleces of iron embedded in a fused

mixture of MgO, Al₂O₃ and SiO₂

M Continuous removal of arrangels from the seaction windows.

en K. lor	museum synthesis
1:K)	K,
200	717 × 40°
300	2.59 × 10 ⁴
400	3.94 ≠ 10°
500	1.72 = 10 ³
600	4.53 = 10
700	2.96 = 101
600	3.96 × 10 ⁻¹

of Mil, from the reaction minters

The equilibrium minimizer contains 35% NH₂ by volume. The minimize is cooled by tendent colls smill NH₂ liquefies at -33.0°C and in removed, while N₂ and H₃ remain to the because they are not liquified at this temperature due to loss boiling points. Unreacted the Hallet specified.

About 13% of nitrogen figation on earth is carried out by Heber's rescens.

Haber's process produce 110 milions ions of Nrig in the world.

About 80% of NH₀ is used for melting fertilizers. Some NH₀ is used formation











Male 's of St

75

61

5500

690

160

55

25 13

200

300

400

600

600

Scurbesis of SO

During manufacturing of H_aSO₄ by contact process, SO₂ is oxidised to SO₄.

The reaction is

→ 25O_→ ΔH = -194 kJ/mol 25O_{bal} + O_{bal}

According to Le-Chatelier's principle maximum yield of SO₃ can be obtained following ways

- [I] Removana SO₃ from reaction mixture or adding more reactants.
- (8) Since there is decrease in number of moles from reactants and products. Hence, large in presence favours forward reaction.
- (iii) Since reaction is exothermic, hence decrease in temperature favours forward reaction Termograture and measure are the rate controlling factors for this process

Increasing temperature decreases the yield of SO_{4} .

e.g. Kc at 200 K is 5500 but at 700 K

It is only 13. Hence formation of SO₄ is less.

High pressure can increase the yield of 50s. Actually, amount of Os is increased to favour forward reaction, instead of increasing pressure.

Optimum Conditions for the synthesis of SO,

In actual process following optimum conditions are

- (f) Pressure = 1 atm
- (II) Catalyst = Solid catalyst, $V_{\underline{a}}O_{\underline{a}}$ or finally divided Pt
- (III) Temperature = Initial heating of gases at 650°C then recycled at lower temperature of 400-500°C to increase the yield of SO,
- (le) Continuous addition of O₂
- SO, is dissolved in H₂SO, to form oleum, which is then dijuted with H₂O to get H₂SO.
- H₂SO₂ is called King of Chamicals. It's use gives an idea of the industrial progress of country

TONIC PRODUCT OF PURL WATER

Water is a poor conductor of electricity. Because self-ionization of water is very still

K_c is given by

Charles Part-

 $g_{c} = \frac{[H^{*}]OH^{*}]}{(M_{c}O)} = 1.8 \times 10^{-10}$ $[H_2O]$

Since $1 \frac{d_{10}}{d_{10}}$ of pure water contains 10000. Therefore, conc. of H_1O i.e. H_2O = 1000 _{18 =} <u>55.55 moVdm².</u>

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Since HaD is present in large excess, therefore, its concentration is almost constant.

So,
$$K_C[H_2O] = (H^*)[OHC]$$

or
$$1.8 \times 10^{-34} \times 55.55 = [H^*][OH]$$

et:
$$1.01 \times 10^{-14} = K_{-} = [H^{\circ}][OH^{-}]$$

Where $K_{\Gamma}[H_{Z}Q] = K_{w} = ionization constant or local product of water$

Tigs, the ionic product of water (K_) at 25°C is

$$K_w = (H^*)[OHT] = 10^{-14}$$
 _____(1,

est of temperature on the value of Ko

waler increases with increase in temperature. The Ku value increses almost 75 times when temperature is raised from 0°C to 100°C. However, increase in value of K. is not regular.

This value shows that if 55.5 moles of pure water are present b dig², then only 10 ⁻¹ moles of water are <u>jonized</u>. It shows for water to very weak electrolyte.

Since H₂O gives equal moles of H⁺ and OH uons.

Hence in neutral H₂O

$$[H^{\bullet}] = [OH]$$

Thus eq (1) can give

$$[H^{+}][H^{+}] = 10^{-10}$$

$$[H^*]^3 = 10^{-14}$$

$$[H^*] = 10^{-7} \mod e/dm^3$$

IQH:I(OH:1=10 14

Thus in nautral water, quantity of H*1 and IOH*1 one is always same.





Cy

Colone Charles President

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Chinada Francis

If some and or base is added to water then Key remains same but come of He and Origins is changed.

If an acid is added then

(H0) < ("H)

And if a base is added then

(OH1) > (H1)

Exercise Q12 (a) Define pH and pOH. How are they related with pK_w

pH and pOF

 $\cdot H$

In 1909. Soversom proposed a pH scale, in order to better exorese the concentration of the none in eq. solution.

phi is defined as

The regative logarithm of H* tens cone, is called pH.

Mathematically pH = log [H*]

ce The logarithm of reciprocal of H tank conc. to called pH.

Mathematically $pH = \log \frac{1}{|H^*|}$

or pH = log 1 log [H*]

 $pH = 0 - \log [H^*]$

 $pH = -\log [H^+]$

,OH

pOH can be defined as

The negative logarithm of Off tone cone, is called pOH.

Mathematically pOH = log [OH]

or The logarithm of reciprocal of OH "tone sone, is called pOH

Mathematically $pOH = log \frac{1}{\{OH''\}}$

Relationship between pH, pOH and pKa

Since [H*] (OH) = K, Taking log on both sides

 $\log (|H^*, |OFF|) = \log K_*$

 $\log [H^+] + \log [OH7 = \log K_{\bullet}]$

Multiply throughout by -ve sign

Charles Par-

- log (H*) - log (OH*) = - log K_u

 $(-\log \left(H^*\right)) + (-\log \left(O(H^*)\right) \circ (-\log K_*)$

 $pH + pOH = pK_{a}$

Since pK_e = 14 at 25°C, therefore

pH + pOH = 14 at 25°C

 $pK_w = -\log K_w$

 $pK_{w} \approx -\log 10^{-14}$

pK_ = 14 log 10

pK_ = 14 at 25 °C

Value of pK_ is jess than 14 at higher temperatures.

Table. Relationship of [11,0]; [O3], pH and pCr()				
			is to Artis, part and	hend
	41.0 7	111	TOH :	ptH
P. Oak	7 X 7Q-14	14.0	1 × 10 ⁻⁴	0.0
-37	4 × 10 ⁻⁴	13.0	1 × 10	L.O
. 1	1×10^{-13}	12.0	1 × 10-4	2.0
Biller life	1 × 10 ⁻¹⁷	10	1 × 10 ⁻¹	3.0
messales.	1 × 10 ⁻¹⁰	10.0	1 × 10 ⁻⁴	4.0
100	L x 10-4	9.0	1 × (0 ⁻⁴	50
1500013	! × 10 ⁻⁶	8.0	1 × 10 **	6.0
Nemotral	2 × 10 ⁻⁷	7.0	1 × 10-7	70
ation Ye	1 × 10 ⁻⁴	6.0	1 × 10 ⁻⁴	8.0
	1 × 10 ⁻⁴	5.0	E ≈ 10 ⁻⁴	9.0
A. A. Salar	1 × 10 1	4.0	1 × 10	10.0
Atchesic	, N 10-3	3.0	1 × 10-11	11.0
	1 × 10 ⁻³	20] = 10 ⁻¹¹	+5.0
	1 × 10 ⁻¹	1.0	1 x 10 ⁻¹⁹	13.0
17.124277	1 × 10	ĽΟ	1 x 10 ⁻¹⁶	14.0



Extreles Q12 (b):

What happens to the acidic and lossic properties of aqueous solutions when all varies from 0 to 14.

phi scale generally ranges from 0 to 14.

However, solution with peantive pit and pit greater than 14 are also known.

Generally, lesser the pH, more actile is the solution and vice versa.

Q pid=2 solution is neutral

© PH ≤ 7 solutions are <u>ecidic</u>

Q PH > 7 solutions are basis

College Chemistry: Part-I

Table Approve	nsate pl	Land gOl	i of some common mate	rials of 25°	'C
Platerial Sec.	pH	POH	Material vige	pH	POH
I.O MI HCI	0.1	139	Bread will	5.5	8,5
OU MINCE	1.1	129	Polaties	58	82
#J N CH,COOK	29	J1 10	Rain water 4.	6.2	7.8
Santric palem ger	2.0	.2.00	Milk Com	6.5	7.5
le Lemon .	2.3	11.7	5alina -	6.5-6.9	7.5.7
War View	2.8	.1.2	e Pare water 4	7.0	7.00
J. Solt drinks	3.0	11.00	Eggs (7.6	6.2
Apples	31	10.9	0.1 M.NaliCO.	8.4	5,6
Guardinit	3.1	109	Semuter .	8.5	5.5
Cranges	35	10.5	Milk of magnesia	10.5	3.5
Tomaters A	42	9.8	ec. 0.1 M NH,	11 1	29
A Comment	3.6	10.4	## M Na,CQ,	11.6	24
Bonner .	4.6	94	O.I WINAOH	13.6	1.00

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IONIZATION CONSTANT OF ACIOS (Ra.)

hears some in water either partially or cosmoletely. Marry acids are weak electrosists and they are not 100% tonged.

The average of acids can be ecovered in terms of Kar, which can be defined as follow Consider the ionization of an acid -

Ke in given by

$$K_C = \frac{(H_3 O^* HA)!}{(HA)!(H_2 O)!}$$

Since H₂O is in large excess, therefore its conc. is constant. Hence

$$K_{C}[H_{2}O] = \frac{[H_{1}O][A]}{[HA]}$$

$$K_{i} = \frac{[H_{0}O^{*}](A^{*})}{(HA)} = -(1)$$

Ministration States to the contract of

Where
$$K_{C}[H_{2}O] \simeq K_{a} \simeq \text{tonization constant of acid}$$

Table (5.7) Discontation constants of some acids of

Acid	Dissectation		-
	HQ	K,	Relative strength
HCI		Very large 0 *	Very strong
HNO	HNO ₃ H*+NO ₃	Very large 10 ⁻¹ h	Viny strong
H ₂ SO ₄	H ₂ SO ₄ == H'+HSO ₄ =	-Eign (0·3)	Very strong
HSO,	H50, - H*+50,2	13 × 19 ⁻⁴	Strong
HF	HE # HIAF	67 π 10→	West
сиссон	CHICOOH = H.+CHICOO-	1.8 × 10 ⁻⁴	Wasaic
H*CO*	H₂CO, ⇒ H*+HOO,	4.4 × 10 ⁻²	Week
H ₂ S	H²2 ⇒ H.+H2-	1.0 × 10~3	West
NH ₄	NH ₄ == H" + NH ₈	5.7 × 10 ⁻¹⁰	Weak
HCO,	HCO, H'+00, L	4.7 × 10-11	West
н₀о	H ₂ O == H°+OH−	1.6 × 10-44	Very weak

collections of ionization constant expression

- a Eq. 1) can be used to <u>calculate K</u> for any acid solution. If <u>nH or [H*]</u> of indution. and midal conc. of acid (HA) dissolved is known.
- 9 Similarly equilibrium concentration of H₂O' and A can be calculated, if initial conc of HA and its K, value is known.
- Strength of acid can be judged

Generally

When

 $K_{\rm u} < 10^{-1}$ acid is <u>weak</u>

 $K_{\rm s} = 1 \ {\rm to} \ 10^{-3}$ ecid is moderately strong $K_{\rm s} > 1$ acid is strong

% lonization of week acid can be calculated by the formula

Amount of acid ionized % tonazation = Amount of acid ionized Amount of acid initially available × 100

Should's Distation Less

It states that

Degree of dissociation of a west electrolyte increases with increase in dilution.

Notifical Property of

Parcentage ignization of weak acid depends upon the extent of dilution



Charles English

Generally % tonization of a weak acid increases with increase in direction of solution (i.e. when molarity is uses).

Examples

When 0.1 mole of acetic acid are dissolved in 1000 cm 2 , of solution, then 1.33 moleculars dissociated out of hundred or 13.3 out of 1000. However, when 0.001 moles of acets acid are dissolved in 1000 cm 2 of solution, then 12.6 molecules are dissociated out of 100_0 126 out of 1000.

Value of K, remains constant at all dilutions at constant temperature

Williamity	To be meaging	[41,40]	ren,coon	К.
10000	1.33	0.001330	0.098670	1 79 × ₁0∺
0.0500	1.89	0.000945	0.049060	L82 x 10 ⁻¹
0.0100	4.17	0.000417	0.009583	1.81 × .0
0.0060	5.86	0.000293	0.004707	1.81 × 10
0.0010	12.60	0.000126	0.000874	1.72 x 10 ⁻⁴

Example 4
What is the percentage landation of a salution and in a solution in which 0.1 moles of 0 is been dissolved i dm^4 of the solution. $Ka = 1.3 \times 10^{-6}$ (Segodia flord, 2011)

Solution

CH,COOH = (mol/dm²)

Equilibrium conc. 0.1 - x = 0.1 [since acid is week] x (mal/dm²)

K, for acedo acid is given by

 $K_a = \frac{(x)(x)}{2A} = 1.8 \times 10^{-8}$

 $R_a = \frac{0.1}{0.1} = 1.8 \times 10^{-6}$ $R^2 = 0.1 (1.8 \times 10^{-6})$ $R^2 = 1.8 \times 10^{-6}$

Taking square root on both sides $\kappa = 1.3 \times 10^{-4} \text{ moles}$ Hisnos (H*) = $x = 1.3 \times 10^{-9}$ rapies

45%

Chamled Equipment

% ionazation =
$$\frac{1.3 \times 10^{-3}}{0.1} \times 100 = \boxed{1.3\%}$$

RENIZATION CONSTANT OF BASES (R.)

Weaks bases are generally molecules or long, which take proton from water and produce

Emple

$$NH_{1(eq)} + H_2O_{(f)} = NH_1^*_{(eq)} + OH_1^*_{(eq)}$$

 $QO_2^{-1}_{(eq)} + H_2O_{(f)} = HCO_3_{(eq)} + OH_4_{(eq)}$

Both NH₃ and CO₆² have different strength of accepting proton. The strength of bases can be expressed in terms of K.

Consider hydrolysis of a general base

Ke is given by

$$K_{C} = \frac{[BH^*][OH^*]}{[B][H_ZO]}$$

Since $H_2\Omega$ is in large success, therefore its conc. is constant

$$K_{C}[H_{2}O] = \frac{[BH^{*}][OHT]}{[B]}$$

 $K_G[H_gO] = K_h = ionization constant of base$ Where

Generally

Stration K. . weaker been and vice verse

Chamled Soulibring

	Table 8.9 K, of some imp	artant bases	
Base	Dissaciation	K _i , Frank	Relative Strength
NaOH	NaOH Na+OH*	Very high	very strong
KDH	KOHK+OH	Very high	VETY CTONG
Ca(OH) ₄	Ca(OH) 2Ca*+2OH	High	Strong
NH ₄ OH	NH,OHNH,*+OH	1.81 × 10	Week
CH ₂ NH ₂ (Methyl amine)	CHAN, CHAN, +OH"	4,38 × 10 ⁻⁴	- Weak
C,H,NH, Antine)	CHUNH CHUNH, +OH	4.7 × 10 ⁻⁴⁰	Very weak

ik, and pk

Values of K, and K, are generally very small numbers. These are usually written in exponential form. These can be converted to whole numbers in terms of pK, and pK.

pKa is defined as the regative logarithm of Ka

i.e. $pK_{\bullet} = -logK_{\bullet}$

pKb is defined as the negative logarithm of K,

 $l.e, pK_{b} = -logK_{b}$

Example

 K_a value of acetic acid is 1.8 \times 10 $^{\circ}$. Hence its pK_a is 4.74

- Lastier pK, . weaker acid and vice versa.
- Larger pK₄, <u>weaker</u> base and vice versa
- If two acids have pK, value difference of 1, then acid with smaller pk, is 10 times stronger than the other if the difference is two, then stronger acid is 100 times stronger than the weaker acid,

Exercise Q 13. (a),

What is Louin, Bronsted idea of acids and bases? Explain conjugate acids and bases

LOWRY BRONSTED ACID AND BASES

According to Lowry-Bronsted concept of acids and bases

- ACID: It is a proton thonor or have tendency to donate proton
- BASE; it is a proton acceptor or have tendency to accept proton.

Whenever a weak acid or a base is dissolved in water, a conjugate acid-base pair is produced. produced.

College Chambian, Port-I

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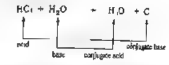
nkei Endlistun

Conjugate Acids and Conjugate Bose

Canjugate acid is a species, which is formed as a result of acceptance of proton by a

je. Every Bronsted acid has a conjugate base. Conjugate base is a species, which is left behind after donation of a proton from the Example:

ionization of HCI in water





Relationship Between Strength of Conjugate Acid Base Pair

Consider the ionization of an acid

Ke is given by

$$K_C = \frac{[H_3O^4][A_1]}{[HA][H_2O]}$$

or $K_{\mathbf{k}}$ is given by

$$K_{\alpha} = \frac{iH^{+}A^{-}J}{(HA)}$$
 ____(1)

Consider hydrolysis of conjugate base

$$A^-_{(aq)} + H_aO_{(0)} \longrightarrow AH_{(aq)} + OH_{(aq)}$$

Base acid acid base

K, is given by

$$K_b = \frac{[HA_[OH\Gamma]]}{[A^*]} \quad [2]$$

Multiply eq(1) and (2)

$$K_a \times K_b = \frac{[H^*][A^*]}{[HA]} \times \frac{[HA][OH^*]}{[A^*]}$$







top Charles (Int.)

micel London

$$K_a \times K_b = K_{--}$$
 (3)

- Eq (3) shows that if K_a for an acid is known than K_b for its comjugate base can be
- Since K_{*} × K_{*} = K_{*} Taking log on both sides

 $\log (K_n | K_n) = \log K_n$

$$\log K_s + \log K_s = \log K_s$$

Multiply throughout by -ve eign

$$-\log K_a - \log K_b = -\log K_a$$

$$\langle -\log K_{\alpha}\rangle + (-\log K_{\alpha}) = (-\log K_{\alpha})$$

$$pK_n+pK_n=pK_n$$

Since pK_e = 14 at 25°C, therefore

Prom eq (3) we have

$$K_0 \times K_0 = K_{\rm in}$$

or
$$K_0 = \frac{K_1}{k_1}$$

or
$$K_0 \alpha \frac{1}{\kappa_0}$$

It shoos that

- Stronger and hove relatively week conjugate base, while
- Week acid have relethens strong continues bear

Chemistry Parts HIMMON ION EFFECT

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The decrease in degree of ionization of a weak electrolyte by the addition of another decirolate having a common ion is called Common ion Effect.

The solubility of a partially soluble electrolyte is decreased by the addition of a more cluble electrolyte having a common ion,

Talk decrease in solubility can be explained by Le-Chatlier's principal.

Precipitation Of KCIO,

KCIO, ionizes in H₂O as

- mm - m

If KCI is added to solution then it rankes as follows

KCl_{ot}
$$\Longrightarrow$$
 K*_{tot} + Cf*_{tot}

Since RCI is more soluble in H₂O than RCIO₆, therefore, it torizes to <u>increase</u> the conc of common lon(K*). This increase, shift the KClO_I dissociation equilibrium in backward direction and KClO_a is precipitated

2 Purfloation of NoCl

For the purification of NeCl, its seturated solution in H₂O is prepared. Through this solution HCI gas is passed

NaCl and HCl lonizes in water as

HCl, due to its more solubility increases cond of common Cl ions in the solution, which shift the NaCl equilibrium to left and NaCl is precipitated.

The addition of a common ton to the solution of a less soluble electrolide automesses Contention and the cone of unionized species increases, which come out a Discipliate.

2. Bali Amalyola

Common ion effect is greatly used in groups of qualitative salt analysis

Ground I

H_eS gas tonizes as

in Group II analysis, the concentration of St ions is reduced by adding HCI to the faiution which is more soluble.



Charlest Earlies

Chambers Period HCI foreizes as

Increase in concentration of H* unts shift the H₂S dissociation equilibrium backward direction and thus conc. of S^b in the solution is decreased. Hence, in concentration of S^b ions only precipitate group II basic radicals in salt analysis.

ட போம் 🖽

NH₄OH ronizes as

To provide low concentration of OH* ions, NH₄Cl is added which is more soluble NH₄Cl_{ps} - NH₄*_{led} + CF_{led}

NH4Cl decreases the OH tons cond by suppressing ionization of NH4OH due in common tons effect

A mixture of NH₂Cl and NH₂OH is used as a group reagent for group III radical in Color Action satt analysis.

Common con effect is also used in <u>qualitative analysis</u> and the preparation of <u>buffer</u>

Exercise Q17 (a) What are buffer solutions? Why do use need them in daily life?

BUFFER SOLUTIONS

A solution that resists in pH changes when small amount of an acid or a boss is added It is called buffer

Properties of Bullet

- 63 It has <u>definite</u> pH
- (ii) its pit is not changed on dilution
- its pH is not affected by time. (iii)
- (tv) Its pH is only <u>slightly chanced</u> when <u>strong</u> acids or basis are added to it.

Importance of butter (Need for a Butter)

- Human blood is buffered at pH 7.35. If this pH goes to 7 or 8, a person may dis
- Some reactions require specific pH. Such reactions can be carried out in a specific pH. specific pH
- Buffers are important in chemistry aiki in many other fields such as molecular hills microbiology, cell blology, soil sciences, nutrition, and clinical analysis

Colone Characteristics Parish



Ecercian Q17.

p) How does the mixture of sodium acetate and aceta acid give us the acidic buffer? of Expelain that a mixture of NFLOH and NFLCI gives us the base buffer?

Preparation of Buffer

ABuffer can be prepared by following two methods.

- Acidic buffer is prepared by mixing a weak acid and its salt with strong base It has pH < 7
 - eg. CH₃COOH / CH₃COONa. So, this maxture gives acidic buffer
- ii) Basic buller is prepared by mixing a week base and its salt with strong acid It has pH > 7
 - e.g. NH4OH / NH4Cl So, this mixture gives basic buffer.

The property of a buffer to maintain its pill, when on cold or a base is added to it, is called Buffer Action

Builder is an application of common ion effect.

Consider the buffer of CH₂COOH, CH₃COONa.

CHACOOPI is a weak acid and ionizes very small, while CH4COONa is a strong security and it conizes in water to greater extent and provides acetate cons

If decreases the ionization of CH₃COOH due to common CH₃COO on and pH of и шоп <u>Іпстеалес</u>,

CH₂COON₂ CH_aCOO + Na*

The buffer is actually a mixture of CH₃COOH and CH₃COO⁺ ions

jupie 8,10 Etter	t of addition of sceta		acetic acid
quigli (and)ë	double (Dissectation	***
0.16	0.00	,3	2 89
0.10	0.05	0.036	4.44
0.10	0.10	0.018	4.74
0 10	0.15	0.012	4 92

Cleaker the concentration of CH, COOH, lesser the pH of solution.







If acid or H.O' ions are added to it, they react with CH.COO ion to give be CH.COOH. Thus, pH of the solution is not changed. Since acetic acid is a weak acid. ers to remein in unionized form.

Similarly, if a base (OFT) is added, it reacts with H_aO+ ions to give back H_aO. Thus of of solution is not changed.

Similarly, basic buffer like NH₄CI / NH₄OH resists in pH changes when acid or base

-destantes the μ (O) Bullet (O) indecisors Equation

For Acid Buffer

Consider a weak acid HA and its salt with strong base NaA. These tonizes as

K, for weak acid is givên by

$$K_{a} = \frac{[H^{a}][A]}{[HA]}$$
 ____(1)

or
$$[H^*] = \frac{K_n[HA]}{[A^*]}$$
 ____(2)

(A 1 is mainly provided by NaA which is gronger electrolyte than HA. Heart, I means the ionization of HA.

Taking log of eq (2) on both sides

$$\log[H'] = \log \frac{K_a[HA]}{[A']}$$

or
$$log[H] = log K_a + log \frac{[HA]}{[A]}$$

Multiply by negative sign throughout

$$-\log[H^*] = -\log K_a \cdot \log \frac{[HA]}{[A]}$$

$$pH = pK_{\bullet} = log \frac{[HA]}{[A^{*}]} \underbrace{\hspace{1cm}} (3)$$

Stros [A 1 is mainly provided by NaA (salt.), therefore it represents the conc. Thus eq (3) becomes.

$$pH = pK_n - \log \frac{(acid)}{(salt)}$$

gate Henderson's equation for acid buffer solution is

$$pH = pK_n + \log \frac{(sait)}{(acid)} - (4)$$

this equation shows that pH of acid buffer depends upon

- pKa of ecid
- Concentration ratio of saft and acid.

This equation can be used to prepare buffer of definite p.H.

Thus by proper selecting acid and taking proper conc. of salt and acid, buffer of required pitantie prepared.

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Set buffer is prepared if ratio of conc. of selt and and are least equal, in this case pH or he builter is controlled by the pK, of the acid.

Ecomple I

In CH COOH/CH COONs buffer

Then dromined (4)

$$pH = pK_a + log1$$

$$pH = pK_a = 4.74$$

Partyle 2: Similarly for HCOOH/HCOONs buffer, If HCOOH) = [HCOONs]

$$pH = pK_s = 3.78$$

If [GH₈COOH] = 0.1 mol dm $^{-1}$ and [CH₈COONs] = 1.0 mol dm $^{-1}$

$$p[H] = p[K_n + \log \frac{[1]}{[0,1]}]$$

$$pH = 4.74 + 1 = 5.74$$

George 3 ff (CH₃COOH) = 1.0 mol dm⁻⁴ and [CH₄COONs] = 0.1 mol dm⁻⁴

$$pH = pK_n + log \frac{[0\ 1]}{[1.0]}$$

pH = 4.741 = 3.74

in Barlo buffer.

Consider the case of NH₄OH/NH₄CI.

Mi-OH is a solution of NH₂ in water and can be represented as

NH_{Blass}+H_EO_{ds} NH_B⁺sep + OH⁺sep

Solving this eq. we get Henderson's equation for basic buffer

$$pOH = pK_b + lng \frac{(salt)}{(base)} - (5)$$

This equation shows that pOH of basic buffer depends upon

- pK, of base
- Concentration ratio of selt and base

uple 5 white the pH of a buffer volution in which 0 11 motor CH₂COONs and 0.09 m no are present. Ke for CH₂COOH to 1.8× 10⁻⁶ (Lahore Board, 2014)

$$\begin{array}{lll} & \text{(CH_{2}COONs 1)} & = 0.11 \text{ g/dm}^{3} \\ & \text{(CH_{2}COOH 1)} & = 0.09 \text{ g/mol} \\ & \text{K, of acetic acid} & = 1.8 \times 10^{-6} \\ & \text{Thes} & \text{pK}_{s} = -\log K_{s} \times \log (1.8 \times 10^{-6}) = 4.74 \end{array}$$

Hence according to Handerson's eq.

$$pH = pK_a + log \frac{(CH_bCOON_a)}{(CH_bCOOH)}$$

$$pH = 4.74 + \log \frac{[0.11]}{[0.09]}$$

Exercise Q17 (c)

Exploin OR Describe the term buffer capacity. (Feleclobed Board, 2009: Lahore Board, 2011)

B. Her Capacite

The a mability of a buffer solution to maintain definite pH is called buffer capacity. it's, determined by the actual motarities of its compounds

Controls

Conside, * CHaCOOH / CHaCOONs buffer

if $|CH_aCO/OH| = 0.09 \text{ mol dm}^{-6}$ and $|CH_aCOONa| = 0.11 \text{ mol dm}^{-6}$

Then according see Banderson's eq.

of Continue

A STATE OF THE STA

$$pH = pK_n + log \frac{[0.11]}{[0.09]} = 4.83$$

how if acid or base is acided to this builder solution, it will try to maintain its pH. There will be only slight change in p.H.

Suppose 0.01 moles of NaOH is added to 1 dm³ of buffer it provides 0.02 moles of OH jers, which will react with 0.01 moles of H none. Thus, cone of CH₂COCH will be increased and come of CH₂COCH will be increased.

Hence remaining cond of $CH_aCOOH = 0.09 - 0.01 = 0.08 M$

And cone of CH₃COO Na (salt) =
$$0.11 + 0.01 = 0.12$$
 M

Hence pH = pK_a +
$$\log \frac{[0.12]}{[0.09]}$$

$$pH = 4.74 + log \frac{[0.12]}{[0.08]} = 4.93$$

Hence pH of buffer is only stightly changed.

Addition of 0.01 moles per dra of solution will change the pf-1 from 7 to 12 in pure

SCILIBRIA OF SLIGHTLY (SPARINGLY) SOLUBLE IONIC COMPOGNDS: SULLBILITY PRODUCT)

The product of molar concentrations of apposituly charged lone, in equilibrium with is solid salt in the enturated solution, at a given temperature is called solubility y

It is denoted by K.

imples and Explanation

Consider a goarfnoty soluble sait PbCl₃ dissolved in water

According to saw of Mass action.

Since PbCis is spaningly soluble, therefore, its conc. will remain constant.

$$K_{a}[PbC]_{a} = [Pb^{a*}](Cl^{*})^{a}$$

$$K_{ab} = [Pb^{ga}][C]$$
 1s

where K_{ν} [PbCl₂] = $K_{\nu\nu}$ = Solubility Psoduct constant

Consider PhSO₄

According to law of Memercion.

$$K_{s} = \frac{|p_{0}^{-1}|[so_{4}^{2-1}]}{|p_{0}so_{4}|}$$

Since PLSO₆ # <u>spiniouly noisble</u>, therefore, its conc. will <u>perpain constant</u>.

$$\label{eq:Km} K_m = (Pe^{a_m})(SO_a^{-a_m}) = 1.6 \times 10^{-a} \text{ at } 25^{0}C$$

Where K_{ϵ} [PbSO₄] = K_{n_0} = Solubility Product

Similarly for PbCl₂ K₄₀ = [Pb²⁺](Cl⁻]²

Value of K., is termerature dependent.

Generally K., for a marindy soluble selt is the provinct of the ions, each raised to a resource equal to the co-efficient in the balanced chemical eq.

lor a general seaction.

Europia

$$\begin{aligned} & \text{CaF}_{\text{Superpose}} + 2F^*_{\text{loop}} + 2F^*_{\text{loop}} \\ & \text{K}_{\text{tot}} \circ \left[\text{Ca}^{2+} \right] \left[\text{F}^{-} \right]^2 \end{aligned}$$

- $\underline{K_m}$ in the measure of how far to the <u>right dissolution proceeds</u> at equilibrium t^2
- Smaller the K. ... leases the capability to be dissociated

atile K ta	lnes for some music	compounds to	anquamete z	re arranged alpl	afairre ally)
Safe	Indeprendice t	ĸ.	Sale	for product	K
Agtir	[Ag* Br]	5.0 × ±0-12	Cus	(Co2+ 191-)	8 × 10
Ag _i CO ₁	[Agr)*(CO, **]	8.1 × 10 ⁻¹³	Pa6	[Fe*-1[S*-]	6.3 × 10
AgCI	[Ag*][CI*]	1.8 × 10 ⁻¹⁹	Fe ₂ S ₃	[Fa ¹ *][S ² *-]*	1.4 × 10
Agi	[Ag* []**]	8.8 x 10 ⁻¹⁷	Fe(OH) _a	(Felt (OH-F	1.6 × 10
Ag/S	(Ap*19(90-)	8 × 10-44	Hg5	Hg**}(5 ¹	2 × 10
Al(OH) _e	MAN HOH!	3 × 10 ⁻⁴⁴	MgCO _a	(Mg*-HCO,*-1	3.5 × 10
BaCO _b	[Be F][CO,F]	2 × 10 ⁻⁰	Mg(OH) ₂	[Mg*+][O]+-]*	5.3 × 10
DeSO _o	(Be**)[SO,*-]	1.1 × 10 ⁻¹⁰	MrdS	(Mn ³ *)[9 ⁶ -1	3 × 10
CAS	[Cd*:][St-1	8.0 × 10 ⁻⁶⁷	PbCl ₃	[Pb*+](Cl*-) ^X	1.6 × 10
CaCO _B	(Cab((CO ₃ =)	3.3 × 10 ⁻⁴	MCO.	[Pb*+][CrO ₄ *-)	2.9 × 10
CaP _b	[C ₁ 1+](p−)0	3.2 × 10 ⁻⁴¹	PbSO,	P\$**][SO4***]	1.8 × 10
Ca(OH) _p	(Caf+)(OH+)	6.5 × 10 ⁻⁴	PbS	[95**][5**)	8.0 × 10



collections of Salutable Product

(You may write Example 6 in

LONG CLESTICK OF ON exemple

The state of the s

of Q15 (b) above)

refer Q18 (b)

How do you determine the solubility product of a subgard 100 g of water?

Betermination of Kap from solubility

- If the solubility of a salt is known then is K_{μ} value can be calculated by using $K_{\mu\nu}$
- Solubilities are given in number of <u>seams of solubility</u> of H₂O.

 Since density of H₂O is 1 g cm⁻¹, therefore, solubility can be written as number of <u>seams</u>. of solute/100 cm of solution.
- From this amount of solute / dm^b is calculated.
- Then emount in grams is converted into moles.
- Using <u>balanced</u> chemical equation, K_{er} can be calculated

Company of the second of the s

The solidality of PhF₂ at 28°C to 0.64 gider² Calculate K_{ee} of PhF₂ (Softmania Board, 2011 Molton Board, 2011, Bargadha Baard, 2013)

solubility of $PbF_{\pi} = 0.64$ grdm⁸

i.e. Moss of PbF₂ = 0.64 g Moi. Mess of PbF₂ = 245.2 g mol⁻¹ Thus no. of moles = $\frac{0.64}{245.2}$ = 2.6 x Mi⁻¹ moles

^{According} to Balanced chemical ed.

2.6×10-8 2×2.6×10-8 Mar schooling Wagner, mp(king) 0

Hence $K_{th} = [Pb^{t+}][F^{-t}]$

 $K_{\rm ep} = [2.6 \times 10^{-8}] (2 \times 2.6 \times 10^{-8})^{-8}$

Km = 7 × 10-3







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IYou may write Example 7 in LONG QUESTION as an example

of <u>Q18 (c)</u> above)

Q16 (c)

Hore do you calculate the solubility of a substance from the value of solubility product

2. Determination of Solubility from Kgr

For this, formula of compound and Kip must be known. Unknown molar solubilly S can be calculated using balanced chemical equation.

Table: Relationship between K _a and the solubility of sume compounds.							
Formula (5)	No. ed ions	Cation Amon	Ŕ ₄ ,	Solubility gdm			
MgCO ₁	2	1/1	3.6. × .0 ⁻⁴	19× 10-			
Pt60,	2	1/1	1.69 x 10 ⁻⁴	13 x 10			
BaCrO ₄	2	1/1	196 × 30-10	14 x 10			
CalOHIy	3	1,/2	6.5 × 10	1,175× 10 ⁻²			
SaF _g	3	1,/2	1.35 × 10 ⁻⁴	7.2 × 10 -			
CaF _g	3	1/2	3.2 × 10-11	2.0 x 10 ⁻⁴			
Ag ₄ CrO ₄	- 3	2/1	2.6 × 10 ⁻¹³	8.7 × 10-5			

Example 7

Ca(OH), is a sparingly voluble compound, its volubility product in 6.5 \times 10 4 . Calculate the solubility of Ca(OH), (Garancolo Board, 2008: Balcanipus Board, 2011 Labore Board, 2006)

 K_{sp} of $Ca(OH)_z = 6.5 \times 10^{-4}$

We know موحد اعتقما

Ca(OH)_{2(an)}-Ca24 (4) + 20H (40) Ca(OH) 0 2S Ca(OH)24

Al equilibriu Hence

 $K_{\theta\theta} = \{Ce^{\theta\theta}\}[OH^*]^{\theta}$

 $K_{av} = [S][2S]^2 = 6.5 \times 10^{-6}$

 $4(S)^3 = 6.5 \times 10^{-4}$

 $[S] = \sqrt{\frac{6.5 \times 10^{-6}}{4}}$

 $|S| = 1.175 \times 10^{-2} \text{ moVdm}^2$

Hence at equilibrium

 $[Ca^{2+}] = 1.175 \times 10^{-8} \text{ mov/dm}^6$ and $[OH] = 2 \times 1.175 \times 10^{-2} \text{ mol/dm}^6$

Charleton Ports

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Since

1 mole of Ca(OH), gives 1 mole of Ca³⁺ ions, hence

Solubility of Ca(OH), $= |Ca^{2}| = 1.175 \times 10^{-2} \text{meV/dms}^{\circ}$

4. Effect of common ion on solubility

Presence of a common ion decreases the solubility of a montroly soluble salt

Consider saturated solution of PoCrO.

 $PbCrO_{4j=0}$ $Pb^{24}_{(ad)} + CiO_4^{-24}_{(ad)}$

Now if NaCrO₄ is added then

Na₄GrO_{44eq} ====2Na⁺_(eq) + CrO₄ + _(eq)

Since Na₂CrO₄ is more soluble and provide common ion CrO₄*-Therefore, equilibrium of PbCrO_a goes backward and its solubility is decreased.

Calle	Chambro Parki		Mind Frank
	ORTHODAL AND SHORT AND	att R. OUI SHOAS (Lve)	iser
(4)	Multiple choice questions For which system does the equilibrium co (a) $N_2 + 3H_2$ so $2NH_3$ (b) $H_2 + I_2$ so $2H_3$ (c) $2NO_3$ so N_3O_4 (d) $2HF$ so $H_3 + F_2$ other flowed, 2012 (C.C. When Secret, 3012) (Supremote bosons)		raceminatic
	Which statement about the following equi		
	2SO ₂ + O ₈ = ± 2SO ₂ (a) The value of K _p falls with a rise in Ten (b) The value of K _p falls with increasing P (c) Adding V _F O ₂ catalyst increases the equilib (d) The value of K _p is equal to K _c (a) 2012 (Labora boxed, 2015)	AH = -188.3 kJ/mol sperature ressure	
(880)	The pH of 10 ⁻³ mol/dm ² of an aqueous so	fution of H.SO, in	
	a) 3 (b) 27 (c) 2	(d) 1.5	
False	dabad Board, 2009) (D.O. Khon Board, 2012) 2018 (Laborr I	oard, 2014)	
(False	The solubility product of AgCl is 2×10^{-4} of Ag* ions in the estation is (a) 2×10^{-19} mol dm ⁻² (b) 1.41×10^{-4} mol dm ⁻³ (c) 1×10^{-24} mol dm ⁻⁴ (d) 4×10^{-24} mol dm ⁻⁴ (d) 4×10^{-24} mol dm ⁻⁴ mol dm ⁻⁴ (e) 4×10^{-24} mol dm ⁻⁴ (f) 4×10^{-24} mol dm ⁻⁴ (e) 4×10^{-24} mol dm ⁻⁴ (f)	rd, 2014)	
(+)	An excess of aquaous alliver attracts is	added to aqueous barlu	m chleride
	precipitate is removed by filtration. What all Ag* and NO ₃ -2 only (b) Ag* and Be** and NO ₆ -4 (c) Be ²⁺ and NO ₅ -1 only (d) Be ¹⁺ and NO ₅ -1 and Cf ²⁺ result soort 3017, 2015	are the main lous in the fi	Hrate?

Synthesis of SO₆ is an excitantic propagation of SO₆ is an excitantic probation, measuring to La-Chastlein's panel lacrosse in temperature will shift the procide backward dispersion. Hersali, increase in condne

Substitution of the substi

des Charleton Part I 471 Charles Earlie in H_SO₄ is a strong acid. One motivate of H_SSO₄ has strong acid. One motivate of H_SSO₄ plans two 2^{h^2} ions. Thiswhere 10^{h^2} mod 4^{h^2} m H_SO_4 gives $2 \times 10^{h^2}$ mod 4^{h^2} of 10^{h^2} ions. Thus $10^{h^2} = 2 \times 10^{h}$ is calculated as $10^{h^2} = 10^{h^2} = 10^{h^2} = 2.7$ Ď $E_{\rm tot} = [40^{\circ}] [C1]$ $E_{\rm tot} = 3 \times 3 \times 3^{\circ} = 2 \times 10^{30}$ $\times = \sqrt{2 \times 10^{-10}}$ x = 1414 × 10⁴ fence, this concentration and dan ⁴ AgNO, and SeCl, are soluble in water When on excess of equeue sites ratrain a actual to equ instrum chloride, them following reaction acoust 24g/NO₂+ BaCl₂ 2g/G + 8g/NO₂+ Ag/G + 8g/NO₂+ Ag/G + 8g/NO₂+ Ag/G + 8g/NO₂+ Ag/G is under insolubble so it is removed from the solution by filtration. Since, Ag/NO₂ is additived and removed as Ag/G. SolvO₂, is under soluble. If instalate in the solution in the form of \$B^{-1}\$ and NO₂. Further, since acousts of Ag/NO₂ is added, therefore, Ag of all also be passent in the solution. Hence, Ag^{*}, 8g^{**} and NO₂, with be present in the solution. Q2. Fill in the blanks at which a reaction proceeds is directly (I) Law of mass action states that the (ii) In an exothermic reversible reaction, _ equilibrium towards the forward direction. (91) The equilibrium constant for the reaction $2O_3$ —— $3O_b$ is 10^{86} at 25^{9} C, it tells that czone is _____at room temperature (iv) in a gas phase reaction, if the number of moles of reactants are equal to the number of moles of the products, K₀ of the reaction is ______ to the K_r (v) Buffer solution is prepared by mixing together a weak base and its salt with _____ or a weak acid and its salt with _____

Answers:

States non-time

(c. strong and strong base

Q3. Label the sentences True or False

and Strong both reactoris and products are

(ii) When a reversible reaction attains equilibrium both reactants and products are present in a reaction mixture.
 (iii) The Ke of the reaction.

Charles September 150

The K_C of the reaction A+B \longrightarrow C+D is given by $K_C = \frac{|C|(D)}{|A|(B)}$



nicol Equipme

Therefore it is assumed that [A] = [B] = [C] = [D]

(iii) A catalyst is a compound, which increases the speed of the reaction and consequently increases the yield of the product.

4.2

consequently increases are years at 25°C is 10° in mol² dm⁻⁶ and is represented by an expression $K_W = [H^*][OH^*] = 10^{-14} \text{ mol}^2 \text{ dm}^{-4}$

(v) AgCl as a spaningly soluble ionic solid in water. Its solution produces excess of Ag* and Cl* ions.

Jej Pals mit Palse fill Palse list Trite of time

(a) Exploin the terms "reversible reaction" and "state of equilibrium" (Labore Board, 2012; Multan Board, 2012; Sargodia Board, 2014)

Reversible Reactions on page 431 and State of Equilibrium on page 432

(b) Define and exploin the law of state action and drive the expression for the equilibrium constant Ke.

Solved on Page 433

College Chambridge Partel

(e) Write $R_{\rm c}$ for the following reactions

(i) $Sn^{2*}_{\rm cut} + 2Fe^{2*}_{\rm cut}$ (ii) $Ag^*_{\rm put} + Fe^{2*}_{\rm cut}$ (iii) $N_{\rm sut} + O_{\rm sut}$ (iii) $N_{\rm sut} + O_{\rm sut}$ (iv) $4NH_{\rm sut} + 5O_{\rm sut}$ (v) $PCl_{\rm sut}$ $PCl_{\rm sut} + Cl_{\rm sut}$

(i) $K_C = \frac{(Sn^{4+})[Fe^{2+}]^2}{(Sn^{2+})[Fe^{2+}]^2}$	(ii) $K_C = \frac{[Ag][Fe^{3a}]}{[Ag^*][Fe^{2a}]}$	(iii) $K_C = \frac{[NO)^2}{[N_2][O_2]}$
(iv) $K_c = \frac{[NO]^4 [H_2O]^6}{[NH_3]^4 [O_2]^6}$	$[v] K_C = \frac{[PCl_3][Cl_2]}{[PCl_8]}$	

Q5. (a) Reversible reactions exicin the position of equilibrium, which is dynamic in natural (Surgodho Board, 2012) d not statte. Explain it,

Off studify that chemical equilibrium is dynamic in nature.

[Fates[abad Board, 2009; D.G. Khan Board, 2009]

At equilibrium stage, the concentration of reactants and products become constants and products become constants. Thus, there may be two possibilities.

* Either all the reactions have stopped in the system and the system becomes stationary, or

both forward and revene reactions are occurring at the same rate.

Second possibility is generally accepted. Hence a chemical equilibrium is called as a dynamic equilibrium.

College Chemistre: Part-

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Character Southers

(b) Why do the rates of forecard reactions slow don able rec approaches the equilibrium stage?
|| |s due to two reasons.

(i) According to law of Mass Action, the rate of reaction is directly proportional to the concentration of reactions. As the reaction proceeds, the concentration of reactions decreases with time, so the rate of forward reaction decreases

(ii) With the passage of time, the concentration of product increases, so the rate of backward reaction also increases and thus rate of forward reaction slows down

At dynamic equilibrium state, rate of forward reaction becomes equal to the rate of backward reaction.

When a graph is plotted between time on X-axis and the concentration of reactants and products on Y-ade for a resmalble reaction, the curve becomes paralisi to time axis at a certain stage,

il At what stage the curves become parallel?

The curve becomes parallel to time axis at equilibrium state. At this state,

it the concentration of reactants and products become constant. So, there will be no change in their concentration with time.

(II) the rate of forward reaction becomes equal to the rate of backward reaction, so the equilibrium is dynamic in nature.

(b) Before the curves become parallel, the steepness of curves falls? Give reasons

In the beginning of reaction, the steepness of curves is more it is because the rate of forward reaction is tast because concentration of reactant is high. With the passage of time. the concentration of reactants decreases, thus the overall tate of reaction decreases, hence steepness of curves fall until equilibrium is reached.

(c) The cole of decrease of concentrations of the mechanic and rais of increase of compensations of any of the products may or may not be equal for various types of reactions before the equilibrium time. Explain it.

Consider the following reaction

2A + B + 3C

This reactions shows when 2 moles of A seads with 1 mole of 8, then 3 moles of C are Roduced. Thus, rate of decrease in concentration of A is twice the rate of decrease in concentration of B. Similarly, the rate of immediate of B. Simmediate of B. Similarly, the rate of immediate of B. Similarly, ontogenhation of B

So tale of decrease in concentration of reactant or rate of increase in concentration of Product depends upon the nature of reaction.

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Chamical Postings

College Charles on Paris

47.1

(a) Write down the relationship of different types of equilibrium nonatents, i.e. it and No for the following general reactions:

Solved on Page 439

(b) Decide the comparation magnitudes of Kg, K, for the following reactions.

Sauthoris of NH;

Ke is given by

K, is given by

$$K_P = \frac{P_{NeG}^3}{P_{N_b} \times P_{H2}^3}$$

For this reaction, change in number of mokes is given by \$\Delta = \text{sumber of mokes of product - number of mokes of reactants}

=2-(1+3)=-2

$$\mu_p = \mu_n \times (RT)^{-4}$$

$$cor \quad K_{\sigma} = K_{\alpha} \times \frac{1}{(RT)^2}$$

Thus if T is such that RT > 1, then $R_{\nu} < K_{0}$

If T is such that RT < 1, then Ky > Kc

No is given by according to law of Mass Action

N_e in given by

For this reaction, change in mumber of moles is given by

All - marker of coules of product - marker of product of sectoris *2-1-1

$$\hat{H}_{\alpha}=\hat{g}_{\alpha}\circ(\widehat{B}T)$$

These BY as such that RY > E_i then $E_p > E_R$ BY a such that RY < E_i then $E_p < E_R$

Charleton Ports



(a) Write down K_0 for the following reversible reactions. Suppose that the solution of reaction mixture in all the cases is V' and at equilibrium stage.

(i) CH,COOH+ CH,CH,OH = CH,COOC,H,+ H,O

(ii) H₈ + I₂ = 2H(1)
(iii) 2H(1 = H₃ + I₂
(iv) PCI₃ = PCI₅ + CI₆
(v) N₆ + 3H₃ = 2NH₆

(b) How do you explain that some of the reactions mentioned above are affected by okunge of volume at equilibrium stage?

(1) The reactions in solution phase are not affected by volume changes, since, in solution $\Delta V = 0$ 2) The reactions in gas phase are of two types.

(i) the reactions in which number of makes of mactants are equal to the number of moles of products. Such reactions are not affected by volume changes.

 $2HI K_c = \frac{\pi_A}{(a-x)(b-x)}$ Example. Ha + Ia 😑

The K_0 expression for this reaction does not involve volume (V) terms, each is not

affected by volume changes (or pressure changes).

(ii) the reactions in which number of moles of reactable are not equal to the number of moles of products. Such, reactions are affected by volume changes.

x2 Example: $PCl_0 \Rightarrow PCl_0 + Cl_0$ $K_c = \frac{A}{(a-x)V}$

The K_{ϵ} expression for this reaction involves, Ψ term so, it will be affected by volume changes (or pressure changes)

99. Explain the following two applications of equilibrium constant. Give enamed (i) Direction of reaction (ii) Extent of reaction

Solved on Page 441 (Direction of Reaction) and Page 442 (Extent of Reaction)

Q10. Biquichs the joilnesing with recents.

b) The change of column disturbs the equitorium position for semi-Perceions, but not the squalleries constant, franched lived, 200, 12m front, 2001.

The reactions is gas phase one of two types.

the tractions to which number of moles of markets are squal to the number of moles of

products, Such, emotions are not affected by solution changes.

Stemple: Fig. + 1,

2.22

The marriage in which parameter of motor of specimens are not equal to the marriage of motors of specimens in which parameter of motors of specimens. Make of products. Such reactions are affected by volume changes.







Annual Company of Particle

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Charles Employee

Promptix $PCI_k \neq PCI_k + CI_k X_1 = \frac{A}{(d-x)V}$

The K, expression for this struction trevolves, V term so, if pressure is increased then The K, expression for the section the reaction will proceed in backward direction to keep according to Le-Chatter's principle, the reaction will proceed in backward direction to keep according to Le-Unemer's presuper, an additional position will change but K_C will remain the constant value of K_C. Hence, equilibrium position will change but K_C will remain constant.

(b) The change of compercious disturbs both the equilibrium position and the equilibrium (Surgoine Board, 2010; Lohors Board, 20.1. Multan Board, 2012) monature of a reaction.

The temperature change affects the anthaipy of the system. Thus, if temperature is shanged, the reaction will proceed either in forward or backward direction. As a rest concentration of reactants and products will change and thus value of Kc will also be changed. Hence, both equilibrium position and equilibrium constant (K_c) will be changed.

Example: N_c + 3N_c = 2NN_c

(e) The anhability of glacons is under it increased by increasing the immerciality.
(2.G. Herr Stard, 2002. Columnia Send, 2009. Lahura Board, 2010, 2012: Feinkabed Soyd, 2011.
Inhability Stard, 2012: D.G. Khen Beard, 2012)

The solubility of glucose in water is an endothermic process

Change + H₂O = Change, eq: AH = +ve So, ecceeding to La-Chadless Principle, if temperature is increased, the system will now in forward direction. Thus, more glucose will be dissolved. Hereon, solubility of glucose with increase in functionalists.

(a) What is built product of owner? How does this units may said the cheef b IP is it tens that this value is 15 these often the T of more becomes from 0°C to 100°C.

lanc product of water is given by the equation

Value of H₂ streament with increase in hospositions. It is because increase in represent the streament, the streament of H₂O. Thus, more H* or OH? - stre of reduced blooms about the deced Home where of the learn

たおで X₉=1×10⁻¹⁶, and み加作 X₉=75×10⁻¹⁶

左がた(先)。 - 81 × 30 ⁽¹⁾ た30で(第₁1₁₁ - 75 × 30 ⁽²⁾

Death ag (2) by ag (1)

Charleton Part

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Chieslant Smitheli

STATE OF THE PARTY AND THE

 $\frac{\left(K_{W}\right)_{00}}{\left(K_{W}\right)_{0}} = \frac{7.5 \times 10^{-14}}{0.1 \times 10^{-14}} = 75$

or $(K_{\rm st})_{100} = 75 \times (K_{\rm st})_{\rm s}$ Hance, $K_{\rm st}$ et 100°C is 75 times more than at 0°C

(a) What is the Justification for the increase of looks product with temperature?

(Bahasabur 2008: Rawalpind Board, 2013) OR Why the K_a of under increases with increase to temperature.

The value of Ke increases with increase in temperature. It is because increase in temperature increases the lonization of H₂O. Thus, more H* or OH * tors are produced. Hence value of Ken Increases.

(a) How do you prove that at 25°C to 1 dm² of water, there are 10° motes of H₂O°?

OR How would you prove that at 25°C, 1 dm² of water contains 10° moins of H₂O° and 10° moins of OH4

At 25°C

 $K_W=[H_0O^*][OH^*]=10^{-44}$ __ (1) Since ionization of water gives equal number of H_0O^* and OH* tons, therefore

 $[H_3O^*] = [OH^*]$

Hence, eq (1) can be written as

 $K_{\rm p} = [H_{\rm g}O^*][H_{\rm g}O^*] = 10^{-14}$ $[H_{\rm g}O^*]^2 = 10^{-14}$

Taking square root on both sides

 $[H_2O^*] = 10^{-3} \text{ moleton}^3$

Hence, at 25°C, water has 10°2 molether of H_eO' inve-

QD. (a) Define pH and pGH. How are they related with pK-

OR Water closure the redutionship between pt L pOH and pto

Softed on Plage 450

What happens to site actific and haste properties of squares and chain pill quality

Jun 6 to 14.

Juliani on Page 451

(i) It is true that the main of pill, and pill, is shown equal to \$4 or off years. DG Rep Band 2013

Total Want why?

The man of pK, and pK, is equal to 14 only at 25°C short pK, = 14 at 25°C. If the persons of pK, and pK, is equal to 14 only at 25°C short pK, and pK, is expensed. Thus, when of K, is consent.



College Chemistry, Part f

and pK, decreases (since pK, α , $\frac{1}{K_w}$). Thus, at other temperatures, the sum of pK, and pK_h is not equal to pK_W . Since, K_W increases with temperature, therefore, value of pK_{Wh} less than 14 at temperatures higher than $25^{\circ}C$.

(a) What is Loury Bronsted idea of acids and bases? Explain conjugate acids and bases.

Define acid and base by Lowery-Bronsted concept.

(Rawalpradi Board

(b) Acetic acid dissolves in water and gives proton to water, but when dissolves a H_sSO_q, it accepts proton. Discuss the role of nostic acid in both cases.

Aretic acid is a stronger acid than water So, it donates proton to water and aris as in

CH₃COO" + H₃O* CH-COOH + H-O

However, H.SO, is a stronger acid than acetic acid. Therefore, H.SO, donates proto and acts as an acid while acetic acid accepts proton and acts as a base. $H_0SO_0 + CH_2COOH_2$ — HSO_4 - CH_2COOH_2 +

Q14. In the equilibrium PO What is the effect on

PCl_{s(p)}+Cl_{s(p)} △ H=90 kJ/mol

(a) The position of equilibrium

(b)Equilibrium constant? If

(i) Temperature is increased

The reaction is exothermic. Therefore, according to Le-Chather's principle, increase in temperature favours forward reaction. Thus, concentration of PCl₅ will, decrease in that of PCl₅ and Cl₅ will increase Hence, it will disturb both equilibrium position and the publishing concentration. equilibrium constant (Kc) of the reaction

(ii) Volume of the container is decreased.

In this reaction, numbers of moles of reactants are less than that of product Therefore, according to Le-Chattiers principle, decrease in volume favours backers reaction towards less polymer Therefore reaction towards less volume. This will change the equilibrium position but value of ke remains constant.

(ii) Catalyst is added

The catalyst has no effect on the equilibrium composition of the system. Thus, it was affect both equilibrium position and W not affect both equilibrium position and Ke. It simply necesses the rate of forward backward reaction. Thus, it only described backward reaction. Thus, it only decreases the tune to reach the equilibrium state

(lp) Cl. is added

Cl₂ is the product of reaction. Therefore, according to Le-Chatlier's principle, gracing to the Chatlier's principle, favours backward direction. In this case, equilibrium position will be changed by P value of K_C will remains constant value of K_C wil. remains constant.

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1015 Bydithesis of NH₈ by Haber's process is an exothermic reaction Ng + 5Hg = 2NH2 AH=- 92.46 k2/mol

is What should be the possible effect of change of temperature at equilibrium stage? Since reaction is exothermic, hence according to Le-chattiers principle, decrease in appearance favours forward reaction. The optimum temperature is 400°C

(a) How does the change of pressure or volume shifts the equilibrium position of this

gg What will be the effect of change in pressure or arranging synthesis? (Lakore Roard, 2002) Paintabul Board, 2008: Violtan Board, 2010: Gapranusts Board, 2014)

to the change of pressure shifts the equilibrium position of synthesis of ammonia Galore

Since number of moles of products is less than number of moles of reactants. Therefore, scording to be Chattiers principle, increase in pressure decreases the volume and thus ours forward reaction towards less volume. The optimum pressure is 200-300 atm.

What is the role of the catalyst in this reaction?

The satatyst used in this reaction consists of pieces of from embedded in a fused modure $M_{\rm SO}$, ${\rm Al}_{\rm SO}$ and ${\rm SiO}_{\rm S}$. It simply increases the rate of forward and backward reaction ha, it decreases the time to reach the equilibrium state.

id that happens to equilibrium position of this reaction if Mil, is removed from the reaction seems from time to time.

According to Le-Chattiers Principle, by continuous ternoval of NH, from the reaction aire, reaction moves in forward direction.

Sulpharte until is the king of chamicals. It is produced by the burning of SO_{ϵ} to SO_{ϵ} through an exothermic represide process.

This the belonced reversible rever's

250₃₀₀ # O₃₆₀ ____ 25O₆₆

 $\Delta H = -194 \text{ kJ/mol}$

that is the effect of pressure change on this reaction?

Since number of motes of products is less than number of motes of reactants. Therefore, couldn't to Le-Chattlers principle, increase in pressure decreases the volume and thus to forward reaction towards less volume.

plant Chamber Period permittee of 400-500°C is required to be 42.1 (c) Reaction is exechermic but still the to Chimbool Sandboom the stabl of SO₂. Give remove. the shall of SO₂, come resolvermic, hence according to Larchastiers principle, dances, Willow do you determine the solubility product of a substance when its solubility to Since reaction is excurrent, manual the yield of SO₃ is increased. However, at temperature fearcurs forward reaction and the yield of SO₃ is increased. However, at temperature fearcurs forward reaction. provided in gramm/100 g of water? temperature, the rate of formation of SO, is slow. To increase the rate an opge How can you determine K_from solubility? temperature of 400 - 500°C is used. (Fotosishad Board, 2012) Saled on Page 467 Q17. (4) What are buffer solutions? Why do set need them to daily Iffe? Allow do you calculate the solubility of a substance from the naive of solubility Solved on Page 460 isual on Page 468 (b) How does the minture of sodium acetate and acette acid give us the orbits be-IMPORTANT FORMULAS Salved on Page 461 $p_i^* = -\log \{H^{+i}, pOH = -\log \{OH^*, pK_i = -\log K_i, pK_i = -\log K_i, pK_i = -\log K_i\}$ (d) Exploin that a misture of NH₂OH and NH₂Cl gives us the bosic buffer? H + pDH = 14, $pK_a + pK_b = pK_a \cdot K_b = K_a$ (Renedpinel Bourd, 2010, 2011 Multan Board, 2012: Sergocho Board, SB * Amount of substance inputed × 100

Amount of substance infellifty available. Solved on Page 461 First an acid buffer pH = pV_{i_0} + $log \frac{(sati)}{(scid)}$ (d) How do you justify that the greater quantity of CH₂COONs in weetle acid decrease the dissociating power of acetic acid and so the pH increases. and for a basic buffer pOH = $pK_k + \log \frac{[adt]}{[base]}$ on When the concentration of soil is increased in an addit buffer, then the pH of the sale (Linkore Board, 579) incremer. Why? CH, COOH is a weak acid and ionizes very small, while CH, COONa is a # electrolyte and it lonkes in water to greater endent and provides acetate ions. CH4COOH + H4O - CH4COOT + H4O+ CH₄COO + Na* CH₂COON₃ Thus, CH₂COCNa decreases the ionization of CH₂COOH due to common Chi^{OS} and old of solution in common Chi^{OS} ion and pH of solution increases. (e) Explain OR Describe the term buffer capacity. (Falsafobod Board, 2009: Labors Sont B. Solved on Page 464 Q18. (a) What is the solubility product? Derive the solubility product expension apartingly soluble compounds, AgCl, AgcCrO, and PhCly OR Define solubility product. Derive solubility product expression for AgrCrO₁ and psch

(Spragodko Room, A)

Solved on Page 465



College Chemistre: Partel

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Charles Parks

NUMERICAL PROBLEMS (Cvercise)

Q19. Ke value for valume for the following reaction is 0:016 at $520^{\circ}\mathrm{C}$ 2. He collective to the state $H_2 + I_2$ and $H_3 = 0.01$ M, $H_3 = 0.02$ M, T_4 is solved an observation of the state concentration of [HI], [H], and [I] when equilibrium is re-established.

Calumbian	-4		
Sciention	2HI 	<u>-</u> - H₂	+ l _e
	0.00	0.01	0.01

(moister) 0:01 0.01 0.096

0.096 - 2x 0.01 + x 0.01 + x

According to law of mass action

$$\mathcal{H}_{e} = \frac{\left[H_{2}\right]\left[s\right]}{\left[HI\right]^{e}}$$

$$K_c = \frac{(0.01 + x)(0.01 + x)}{(0.096 - 2x)^2} = 0.036$$

$$=\frac{(0.01+x)^2}{(0.096\cdot 2x)^2}=0.016$$

Taking square root on both sides

$$\sqrt{\frac{(0.01 + x)^2}{(0.096 - 2x)^2}} = \sqrt{0.016}$$

$$\frac{(0.01 + x)}{(0.096 - 2x)} = 0.126$$

$$0.01 + x = 0.126 (0.096 - 2x)$$

$$0.01 + x = 0.0121 - 0.252 x$$

$$x + 0.252 x = 0.0121 - 0.01$$

$$1.252 \times 0.0021$$

$$x = \frac{0.0021}{1.252}$$

Charleton Parts

Concentrations when equilibrium is re-established are

Character Conditions

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The equilibrium constant for the reaction between scatte and end ethyl alcohol to 4. A mixture of 3 males of acetic sold and one male of C₂H₂OH to allowed to come to equilibrium. Calculate the amount of ethyl scotate at equilibrium stops in number of moles and grams. Also calculate the museur of reactants left bekind,

According to law of mass action

$$K_{a} = \frac{[CHaCOOCaHa][HaO]}{[CHaCOOH][CaHaOH]}$$

$$\mathbf{x}_{i} = \frac{(\mathbf{x}_{i}(\mathbf{x}))}{\mathbf{x}_{i}(\mathbf{x})} = \mathbf{4}$$

$$K_{c} = \frac{\left[x\right]\left[x\right]}{\left(6 - x\right)\left(1 - x\right)} = 4$$

$$x^2 = 4(3-x)(1-x)$$

$$x^{2} = 4(3 - 3x - x + x^{2})$$

 $x^{2} = 4(3 - 4x + x^{2})$

$$x^2 = 4(3-4x+x^2)$$

$$x^2 = 12 - 16x + 4x^2$$

$$x^{2} = 12 - 16x + 4x^{2}$$

$$12 - 16x + 4x^{2} - x^{2} = 0$$

$$12 - 16x + 4x^2 - x^2 = 0$$

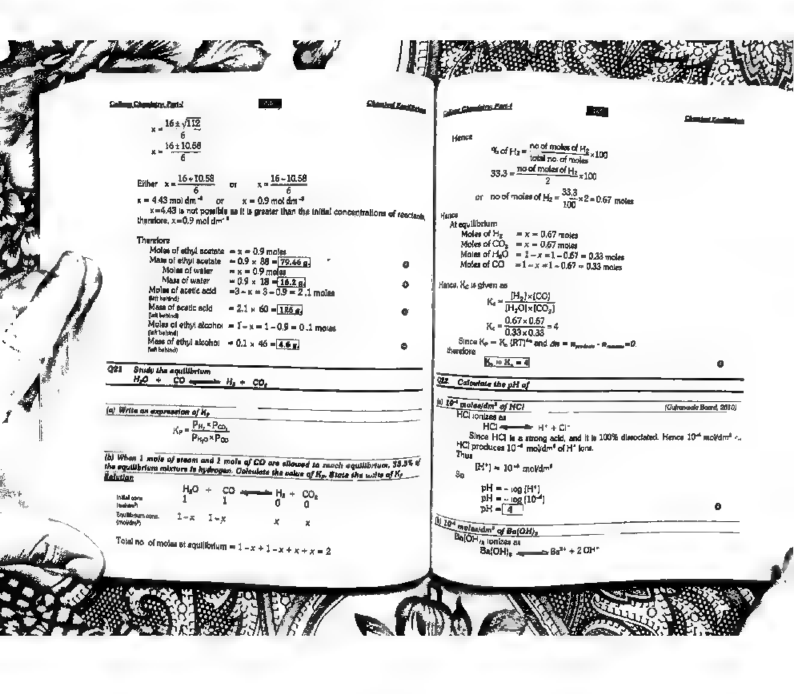
$$3x^{2} - 16x + 12 = 0$$

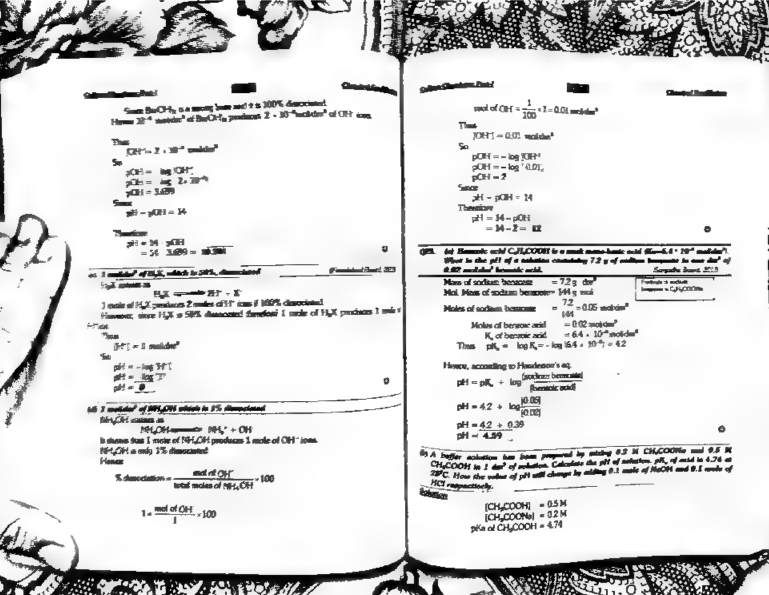
It is a quadratic equation and can be solved by using quadric formula

$$x = \frac{-b \pm \sqrt{b^2 - 4ac}}{2a}$$

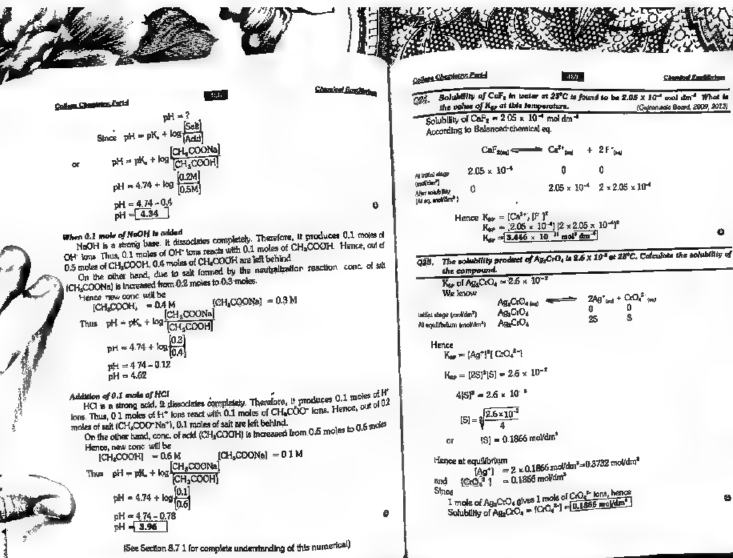
$$x = \frac{-(-16) \pm \sqrt{(-16)^3 - 4(3)(12)}}{2(3)}$$

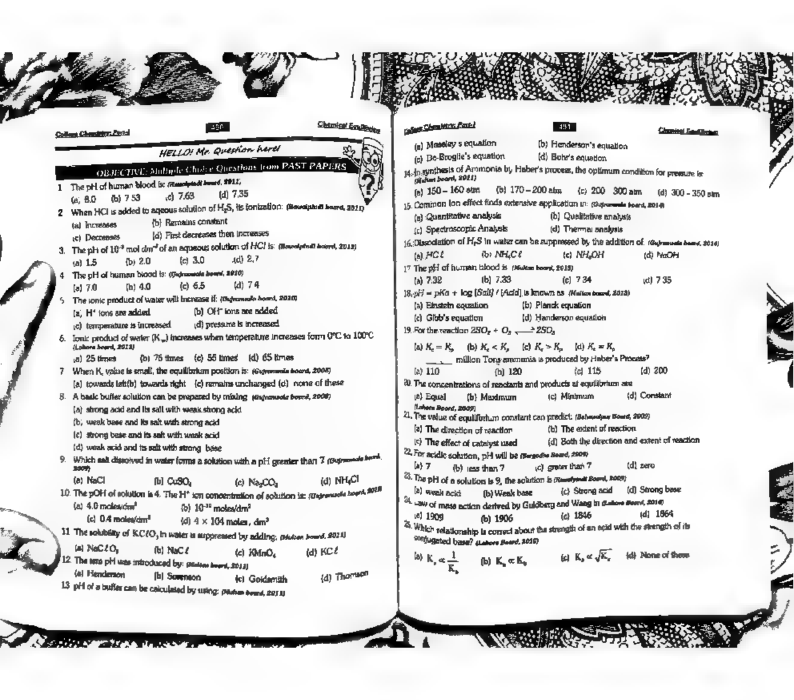
$$x = \frac{16 \pm \sqrt{256 - 144}}{5}$$

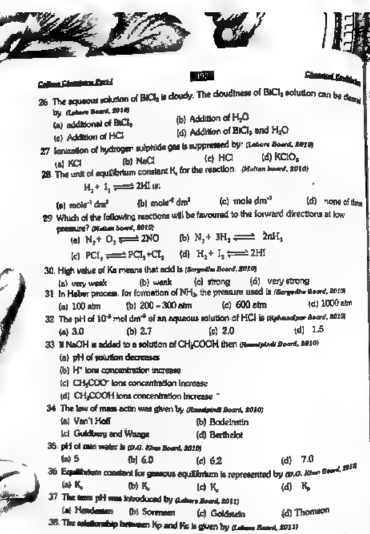




All the







The state of the s

C /5424				THE PARTY	weig g		26 /2	ML CON
San Charles	Parti			49:			Chap	right Employment
(6) K.	≖K _e (P) [™]		(b)	$K_s = K_s \left(\frac{P}{N} \right)$)			
(c) K,	$=K_{\epsilon}(RT)^{44}$			$K_a(RT)^{an}$				
g, The uni	storK, of H, ador³	O are. (Patente (b) moles ^a CH3COONs	dm."	(c) mo	ler ² dm ⁶	ul Ja	(d) n	voles ².dm³
w1.7	(6	:}>7	(0	< 7	{	d)	1	•
(e) 7	(6	OH is always i) Zero	(c	14	- (d) 16	0-14	
(2 n case		> [H*] the soli			oard, 3007)		
(a) Acid		Neutral		Basic			one of I	
(a) Inc. 44 Reactio	eases (to	NH ₄ OH solud) remains ser h H ₂ O gives a no soord, 2011)	16	(c) decreas	ses (d) u	CTV AND	100 times
	ing BICI,			ding HCl				
(c) Iner	easing tempe	rature	(d) de	යැනෝවල කො	perature			

A	Answers to Multiple Choice Questions from Past Papers.									
Q#	Ans	Q#	Ans	Q#	Ans	Q#	Ans	Q#	Ans	
1		47		47.77	(c)	4	(c,	5	(b)	
	(d)	- Z	,c)		(c)	9	(b)	10	(d)	
11	(a)	7	(6)	13	,c)	14	(6)	15	(a)	
	(b)	12 -	(b)	-	(6)	19	(a)	20	(d)	
16	(d)	17	[d)_	18	(6)	24	(d)	25	(a)	
21 26	(d)	22	(b)	23		29	(c)	30	(d)	
40	(c)	27	{c}	28	(d)	34	(c)	35	(a)	
31	_{b}	32	(a)	33	ic -	39	(b)	40	(c)	
36	(d)	37	(6)	38	[3.	44	(b)			
41	(c)	42	(c)	43	(¢)	-1-5	107			

Detailed Explanation of Past Papers MCQs & Qualitiers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK-I



SHORT & LONG QUESTIONS FROM PAST PAPERS

HERR & LAW OF CONSTANT HEAT SUBDIATION, BORN-HABER CYCLE

Short Constinue

Short Quantions: (61 mount Short Countles 1, sto.) Justify that one moles solution of used in water is discher (Behandpur Beard, 2009) Justify that one motor solution of unput in words to close (Sergodhe Read, 2009)

Hoss the solution of copper suiphate is ocidic in nature? (Sergodhe Read, 2009) Mo.SO. 10H,O should discontinuous solubility curvs. Glue reason, (Sergodhe Seard, 2015) Why agreement solution of NH₂Cl is addic but that of CH₂COOH is basic? (Receipted Base), 1918; in the control of NH₂Cl is addic but that of CH₂COOH is basic? (Receipted Base), 1918; in the control of NH₂Cl is addicable that the cont

Calculate pH of 10" mol/dm*of Ba(OH),

Agusous solution of Na₂CO₄ is alimites in notice. Explain, ID.4. Nies Beers, 3007

REVERBILE MEREVERBIBLE REACTIONS, CHRISICAL EQUILIBRAM

Short Questions: (S) means Short Question 1, ste.)

- (1) Define reversible and irreversible reactions. Olive examples. (Asset Kashnir Bord, 2018)
- (2) Differentiate between reversible and integrable reactions. Give examples. (Reads level 3011 D.C. Khan Seers, 2018) (3) Define state of chemical equilibrium. OR what sie you understand by chemical
- equilibrium? (Supments Sourd, 2000; D.G. (Une Sourd, 2012)

LAW OF HASS ACTION

- Short Characterist.

 () Define loss of most action. Studen Search, 2007: D.S. (then Search, 2012: Surpodie S.). (2) Why the aquilibrium constant value has its units for some of the reversible reaction, its
- has no units for some other resultions? (D.G. Khen Sound, 2008; Resemb
- Operation 1) Define law of mass actions. Derive equilibrium constant expression for a general clamb Law Owner raction. (Response Board, 1012; Hulton Board, 2012)

SQUILIBERTH CONSTANT EXPRESSIONS

- (s. Derius K, expression for the reaction, (Labor Sound, 2008)
- CH_COOH + C_H_OH = CH_COOC_H_1 + H_O

 Decree equilibrium communic expression for the dissociation of PCL
- (3) Write aguilliritem area test expression for the reaction. - Avent, Marm con What are 160 and 16 N. + 3H, # 2504,
- 细度 te deu m relation between K, and K, (m.e. exand how these are related? page -t- (h--), 2423)

APPENDATION OF EQUILIBRIES CONSTANT

Colonia Characteric Product

115

- (i) How Ke predict the extent of a chemical reaction? Provided Stand 2011: its Write two applications of equilibrium constant (age-
- redo Barril, 2023. La
- d) riou does the equilibrium constant of a reaction tell as the direction of a charmeal reaction (D.C. Main Board, 2019; Resolved Seed, 2019; Oil how the direction of a charmed reaction (D.C. Main Board, 2019; Resolved Seed, 2019; Oil How the voice of Kc of a reaction lefts to predict the direction of a reventible reaction? (Buyesha Board, 2007) disjunction (Barrel, 2014)

IL CHATLEE'S PRINCIPLE

- Sect Overstons

 II) Gue statement of for Define, Le Chattler's principle (0.0. Rhes Board, 1907; Lehma Board. 1906 Salesment of for Engine Lo Craner a principle (a.c. rain some 1997) June 1909, 2013; Aud Restrate Sound, 2011) OR Defins four of mass action and La-Chattier's principle. (Housefold Board, 2018) And For Lau of Nam Action, See 1999
- (i) In some reversible reactions, the direction of reaction is changed by change in pressure. Glue reason. (Belie alpur Board, 2009
- B) How does a catalyst offect a reversible reaction? (Paintained Board, 1908, 2019). Philam Bo. 1904: Empodus Beard, 2013) OR Describe for What is the) effect of points on equilibrium.

 Baker Board, 2007: Behmadpur Board, 2010: D.O. Khen Board, 2012: Fateshind Board, 2012: Sehmetour Board, 2018) GR A catalyst does not affect the equilibrium constant. Comment 07 K. (Gutromanda Secret, 2006)
- (ii) What will be the effect on the position of equilibrium on the following system (f. (a) temperature is increased, (b) chlorine is added (Lakere Board, 2018) AH = 90 Milmot

PCI₁ = PCI₂ + CI₂

ter Covernos 1) Define Le-Chatellier's principle, Discuss effect of fit change in volume (ii) change in imperature on the formation of ammonia, (0.0, 10m seers, 2015)

TUCATIONS OF LE-CHATLIES'S PRINCIPLE

- est <u>Desilions</u> Il. How amenousla is synthesised by Haber's process. Also give the optimism conditions for
- Promotion (Serpodia Sourd, 2013)

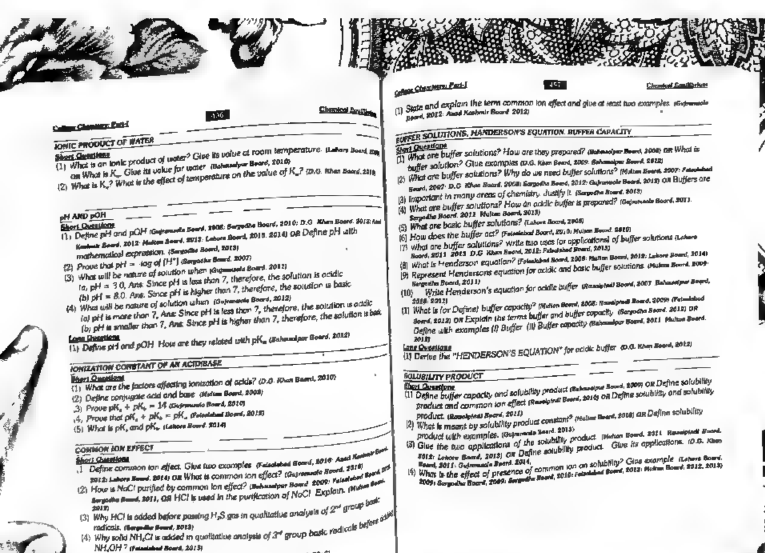
 Transaction of NPI is in Experience of loss temperature. Explain. (D.G. Rim. Sourd, 2007) are Why during the synthesis of NH1, temperature is kept low? Kingsowin 24
- Must all the effect of increase of pressure and temperature on the following resistion?

2NH,(g) 4H = .92.46 kJ.sorn (e)+ 3+(e)

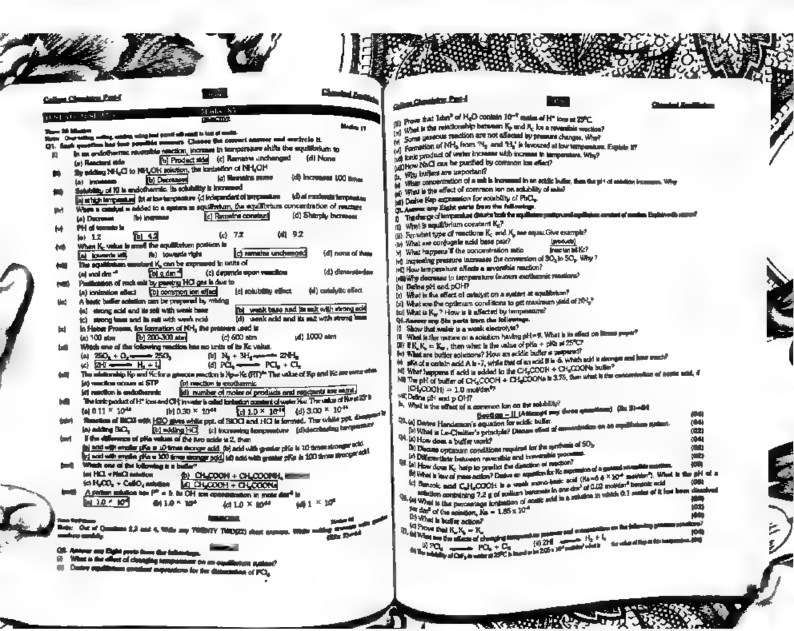
- Total pressure increases the oxidation of SO₂ as SO₃. Explain why? the
- The country pressure increases the customers of the following system of equilibrium materials that will be the effect of volume change on the following system of equilibrium materials that the effect of volume change on the 250s promised forms, 2001 this PCL as PCl₃ + Cl₄ (b) 2SO₃ + O₉ at 2SO₃ photocolomic distributions. See a PCl₃ + Cl₄ (b) 2SO₃ + O₉ at 2SO₃ photocolomic distributions.

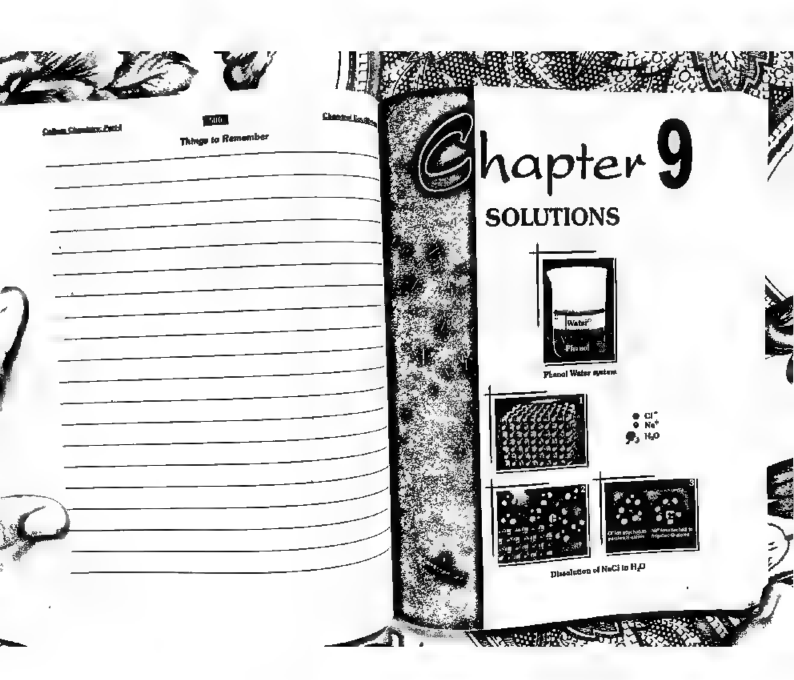
 What a note on synthesis of arranonia gas by Hober's process keeping in mind the Ophicoster of the Polycoping of the Poly

Opheatons of chemical equilibrium in inchary (Labora Beard, 2012)



(5) Give two applications of common ton effect (Outrowells Board, \$6:3)







CONTENTS

Chapter-9

Solutions

CONCEPT OF A SOLUTION CONCENTRATION UNITS OF SOLUTIONS

CONCENTRATION UNITS OF SOLI Percentage composition Molarity (M) Molelly (m) Mole fraction (x) Parts per million (ppm) Interconversion of various concentration units of solutions

representations
TYPES OF SOLUTIONS
Solutions of solids in liquids
Solutions of liquids in liquids
RACULT'S LAW

RAQUEL 5 LAW
Rebuilts law for two volable components
Ideal and non-ideal solutions
VAPOUR PRESSURE OF LIQUID-LIQUID

SOLUTIONS

SOLUBILITY AND SOLUBILITY CURVES

Solubility curves

Fractional crystalization
COLLIGATIVE PROPERTIES OF SOLUTIONS Why some properties are colligative?
Lowering of vapour pressure
Elevation of boiling point
Measurement of elevation of boiling point

Measurement of elevation of busings point
(Lands berger's method)
Depression of freezing point
Measurement of depression in freezing point
Applications of t.p elevation and f.p depression

ENERGETICS OF SOLUTION

Hydration energy of luna HYDRATION AND HYDROLYSIS

Hydration Hydrates

nydrams
(hydrolysis
Objective and short answer, questions (exercise)
Numerical problems (exercise)
(Past Papers MCQs and Short Onestions

Test your skills

50%



CONCEPT OF A SOLUTION

Castr

gile of matter with uniform properties and fixed composition is called a Phase. Exteriple

- Water at normal temperature and pressure is a single phase
- A solution of sugar in water is also a single phase

Soficiano

mus etileture of two or more expetanges to called a solic A^Mhom

Example

Solution of sugar in water.

Trace Solution

A solution composed of two substances is called himsy solution.

Solution of NaCl in H₂O

i de m

e prosper in larger occurre in a solution is called solvent. Avid

interior

À ministance present la smaller anount in a solution is colled solute.

Bample

In 40% augus solution, sugar is solute while $H_{\rm s}{\rm O}$ is solvent.

Brow Sofutani

aliding relatively until amount of solute dis dresietton er ite solution.

Loccompand Solution

A solution containing reg relatively larger appears of a

Bemple

A 10% sugar solution is more constricted than 5% sugar solution.



College Chemistres Probl

Agreeurs Selution

A solution to which water is soluted in called squeous solution.

E comple

Solution of NaCl in H₂O

organication of Solution

nt of solute present in a given amount of solvent is called concentrat The one

Emperior Q6 (a) What are the concentration units of solutions. Compare motor and as schuttons. (For Comparison wer behan molellity on page 507)

CONCENTRATION UNITS OF SOLUTION

Units used to express the ausuant of matrix in solution are called concentration at

Following concentration units are generally used.

l. Percentage composition 2. Mointly

3. Molailly

4. Mole feedion 5. Parts per miles

Parantane Composition

from the expression in feditioning ways

Parameter Weight (Seight (Se eine)

It is the satisfic of solute dissolved per APP posts by weigh Mark to the sales

Table of solution a 100

September 1

A \$55 (units) sugger subdiges its sunter means that 5 g. of solube and present in \$9.9 military in some Time position company To g of water

Colorina dia paramaga by antight of StaCl, \$2 p of StaCl in Associated in 20 p of staCl.

+2g Man of NaCl

Wass of H₂O = 20 g Mass of Solution = 20 + 2 = 22 g

% by wargin of NaCl = mean of NaCl ×100

कार्यक्ष को स्टोक्संबर

To key wanglet of MaCi = $\frac{2}{22} \times 100 \times \boxed{9.09\%}$

College Chemistre, Part.

505

(f) Percentage weight/column (f. wh)

It is the weight of solute discolved per 100 parts by solute of sol

Exemple

A 10% (w/V) glucose solution in water means that 10 g of glucose are present in 100 cm3 of solution.

In this solution, quantity of solvent is not known exactle.

III) <u>Percentage volumejoolgis (% skel</u>

It is the commer of car' of solute dissolved per 100 g of soluti

Exemple

A 10% (s/w) solution of alcohol in water means that 10 cm³ of alcohol and present in 100 g of solution.

In this solution, total volume of solution is not known exactly

(a) Percentage polymericolame (% sin)

It is the volume of white dissolved per 100 cm² of relation.

This unit is used for solutions of liquids in liquids.

Courses

A 12% solution of alcohol in water means that 12 cm² of alcohol are passent in 100 cm² of solution.

For such solutions, total volume of solution may not be some to the sum of volumes of solute and solute.

delactry (M)

Number of moles of solute disorded to I day of solution to called molestic.

It is denoted by 'M'

Mathematically

M = Number of rootes of solute

Volume of solution or deal

Newson of grades = Malecular mate at solute

Since

Thus

Project der mass of solute

Volume of solution in den

 180 g (1 mole) of glucose are dissolved in H₂O and volume of solution is made up to Exercise

I dm^a by adding more H_aO to prepare IM solution



342 g (1 mole) of sucross are dissolved in water and volume is made up to 1 dm⁴ by adding more H₂O to prepare 1M solution.

Sterilarly

Since 342 g of sucrose have larger volume than 180 g of glucose, therefore, in 1 molar sucrose solution, amount of H₂O is sess than 1 molar glucose solution.

Exact volume of solvent can be determined by noting its density.

Exemple 2: elarity of a equation containing 20.7~g of $K_2{
m CO}_1$ dissolved in 500 ce¹ of Culculate the w the gloss solution.

Mass of
$$K_0OO_2 = w = 20.7 g$$

Moles mass of
$$K_zCO_a = M = 138 \text{ g/mol}$$

Volume of solution $= V = 500 \text{ cm}^a = \frac{500}{1000} = 0.6 \text{ dm}^a$

Molestly is given by

$$motionity = \frac{w}{M} \times \frac{1}{V}$$

 $\frac{20.7}{138} \times \frac{1}{0.5} = \boxed{0.3 \text{ mast/shm}^2 \text{ or } 0.8 \text{ M}}$

Solulies on

Number of makes of volume dissolved in 1 kg. of volcent is called mobility.

It is characted by 'm'

Methemetically

m = Number of moles of solute

Mess of solvent in Kg

Mass of solute Number of moles = Molecular mass of solute

Mass of solute

Thus m = Molecular Mass of solute Mess of solvent in Kg

180 g of glucose when dissolved in 1 kg. 1000g) of water gives 1 molal solution of glucose. Total mass of this solution is 1100 - 200. glucose Total mass of this solution is 1180 g. However, total volume of this solution is 1180 g. However, total volume of this solution. is not known exactly To find the total volume, density of solution is required.

Charleton Parel

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- . For 1 modal sucrose solution, 342g of sucrose are dissolved in 1000 g of $\rm H_2O$
- Modality of a solution is an indirect expression of the ratio of moles of solute to solvent.

Comparison of Molar and Molal Solutions

- 1 motel aqueous solution of any substance is dilute than its 1 moter solution. Since in 1 molal solution quantity of solvent is greater
- The value of motality does not change with temperature but motarity changes. It is because molar solutions are based upon volume, which changes with temperature, hence molarity changes. But motal solutions are based upon mass which is not changed with temperature.

ø

What is the molatity of a solution prepared by dissolutes 5 g of tolurns (C,H, in 250 g of (Salamatour Sound, 2011) benzene.

Mass of tokuenet solute) = w = 5 gMolar mass of $C_7H_0 = M = 92 g/mol$ Molar mass of $C_7H_0=M=92$ g/mol Mass of benzane (solvent) = W=250 g = 0.250 kg Motality = ?

Molality is given by

Molality =
$$\frac{w}{M} \times \frac{1}{W}$$

Motality = $\frac{5}{92} \times \frac{1}{0.250} = 0.217 \text{ mod/kg or } 0.217 \text{ m}$

Male Fraction Inc

It is the ratio of the member of moles of a particular component of the solution to the of number of moles of all the components of the solution.

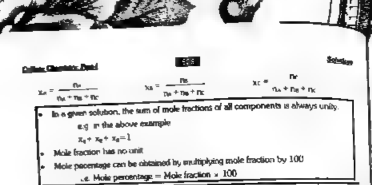
It is represented by 'X'

Mathematically

Number of moles of one component

x = Total number of moles of all components of solution

This unit is also applicable to a solution having more than two components. **G.B. consider three components A, B and C in a solution. Let number of make of A, B and C are n_A , n_B and n_C respectively. The mote fractions of these components in the solution are of the second or the second of these components in the solution are of the second or the s adultion are given by



Calculate the mole fraction and mole percent of each compose g of sales alcohol. 96 g of setting alcohol and 90 g of water.

Mass of ethyl autohol = 92 g Molar mass of ethyl alcohol= 46 g Moles of ethyl alcohol $=\frac{92}{46}=2$ moles

Mass of methyl alcohol = 96 g Molar mass of methyl alcohol = 32 g

Moles of methyl alcohol. $=\frac{96}{32}=3$ moles

Mass of water = 90 g Molar mass of water = 18 g

Moles of water = $\frac{90}{18}$ = 5 moles

Mole fractions are given by

$$x_{\text{ediqui allowhol}} = \frac{2}{2 + 3 + 5} = 0.2$$

$$x_{modest special} = \frac{3}{2+3+5} = 0.3$$

$$\chi_{\text{water}} = \frac{5}{2 + 3 + 5} = 0.5$$

Mole percents are given by

Mole percent of ethyl alcohol = 0.2 × 100 = 20 % Mole percent of methyl alcohol = 0.3 × 100 = 30 % Mole percent of water = 0.5 × 100 = 80 %

9

703

Per Million (ppm)

E to the remainer of parts of solute (by melaid or a light or polarie) of solution.

Methematically

$$pipm = \frac{Mess of solution}{Mess of solution} \times 10^{5}$$

This unit is used to express very low concentrations, e.g. to express the conc. of injurities in water

Comple 5

complete > Sec under > 0.65 > 10 $^{-1}$ g of dissolved acapter in one by of water. Calculate the consequentiation of any general in sea water in prim.

Mass of oxygen = 5.65×10^{-3} g Mass of water = $1 \log = 1000$ g

ppm of oxygen = ?

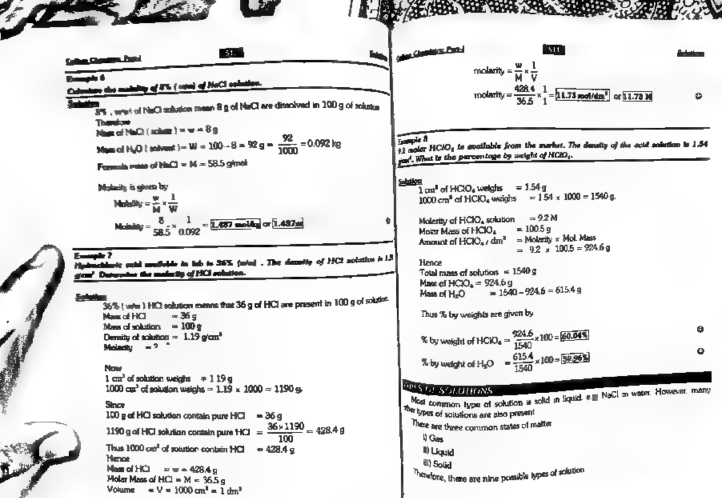
ppms is given by

$$ppin = \frac{5.65 \times 10^{-3}}{1000} \times 10^6 = \frac{5.65 \text{ pain}}{5.65 \text{ pain}}$$

merguension of Various Concentration Unds of Solutions

Various concentration units can be converted into one another. Sometimes chemical vanous concentration units can be converted into one another. Sometimes chemical splits provide solution whose motality is given if we want to know its motality or w/w retentage then we need to convert one unit into other. These conversions are usually done busing formula masses and densities of solutions.

Name of acid	n _e (m or	Malarite (M. dan D	Density Inch. 3
H,30,	98 %	18	1.84
H ₁ PO ₄	85.5 %	4.5	170
	70.4%	15.9	, 42
HNO,	37.2 %	12,1	1 19
HCI -	99,8 %	17.4	1.05



Molarity is given by



Chestone: Parid

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10.5		Table: Com	aini depart and examples of saluther .			
ar.	Solute State	Sylveent Seatte	Company Languages			
	page .	Cent	Air			
<u>.</u> t	Gen	Liepaid	O ₃ its water CO ₆ in water			
- i + (as S			Hi _L adecribed by Pd			
-	Liquid	Gés	Man. Fog. Clouds. Expeld all pollutants			
-	Legita	Jewid	Alcohol in Water mile, bensene in tolume			
-		5old	Marcusy in eliver, butter cheese			
<u>•</u>	Liquid	Gen	Dust particles in smoke			
7/	Solid Solid	Liqued	Sugar in water Jakker, Paints			

Solution of solids in liquids

Dissolution

The selubility of a solid substance in figure is called dissolution

The process of dissolution depends upon forces between solute and solvent.

Example

Dissolution of NaCl in H_gO

Solubility principle states

- " Like dissolves Like "
- ive. Polar substances are dissolved in polar solvents and Non-malar substances are dissolved in non-polar solvents.

In solids, sons or molecules are tightly packed and have strong intermolecular or title prices. The process of dissolution is to break those forces. These forces are broken to solids solvent interactions and these nodes of the collections. by solute-solvent intersections and thus solid is dissolved in the solvent.

In ranic solids, strong exterioric forces are present.

For dissolution, these losses must be broken. Only polar solvents with strong state. ment can break these forces. Hence, tonic solids are woutly dissolved in polar solvents

in non-poise coolecular creates, weak London or dipole-dipole forces are peneral, their to be treated standard by non-polar solvents. Here can be broken assaily by son-polar solvents. Hence, non-polar solids are dissolved in polar solvents. polar scharze.

THE THE PARTY OF T

Chemistry: Part of

terroles.

Conte-sugar with strong hydrogen bonding is poler. It does not dissolve in non-poler solvente such de Herosine oil, petrol, benzene etc. bezauer these solvente can not break 11-bonding of cohe-sugar

But it readily dissolve in H₂O because they develop H-bonding for each other

 NeCl-with strong interioric forces in not dissolved in moderately poles solvent such as acesone. But at once desolved in polar H₂O.

Salution of Liquids in Liquids

It may be divided into three types

Liquide which dissolve together in all proportion are called completely niterible liquide. a Cample who misc thic liquids

Busmole

Solution of atcohol in H₂O

- During formation of such solution volume is decreased or in some cases increases
- Heat may be evolved or absorbed during their turnation
- These solution can be separated by fractional distillation

in Providing arise that liquids

The liquids which dissolve together to some extent are called partially miscible

In other-water system, either (C₂H₂-O-C₂H₃) dissolves using upto 1.2% while maker desolves either upto 6.5%.

Thus it equal volume of other and water are shaken together, two layers are produced

- Water containing ether and
- Ether containing water

Each layer is a saturated solution of other liquid

When two liquids are artised together such that two logers are produced and such layer is a standard politices of other liquid then such solutions are called conjugate solutions.

Note that the conference of th

Musical solubility of liquids in conjugate solutions changes with temperature





the Spinister, statute about the भारता अस स्वीकत ता साम्यान

Thomas Maries session

By streeting injust, violetine of phinoid and water regulating two layers are produced

- "James Aguse an Austral approximations palaconal, hand
- Lower Ager of phonoic continuing with:
- ne policy make more a 5th neutron of observation water, while

the second water it when the

ीर अवस्था केंग्रस प्रतिकार के कारीया प्रतिकार के विकास विद्वार प्रतिकार के विद्यार

िक्षा अन्य कार्याकृतः निकृताः सूत्र असीता कार्यास्त्रकः वर्षास्त्रकः

The same taper has grainer density that to greater processings of physical.

When temperature is increased, more water moves but phonol and more phased and interperature of the minute of phased and water is obtained. The minute contains 4% themselves and 6% years 6% 9% is called supper contains increasing of this system.

The anaparature at which has employed what management and the property of the state of the pro-

Different systems have their count consulate temperature and composition.

Acres 1

Where Arollow system has upper constants temperature of 167 $^{\circ}\mathrm{C}$ with 15% $\mathrm{H}_{\mathrm{s}}\mathrm{O}$

Machinest quintiments spitem has upper consister temperature of \$115 and \$1

contrangulation duministration digitals.

Experiences.

Water - Bensens (1HgO + CgHq)

 CS_k Wester $H_kG + CS_k$

to Charles Park

545

Barrier Q13. What is Repull's law. Give is three statements. How this lose can help us to understand the identity of a solution.

garder's LAW It was given by F.M. Raoult in 1887

posts present of a solvent above a solution, is aqual to the re of para technical and main function of solvent in solution.

Mathematically

$$p = P^{0}x_{k} - \{1\}$$

P = Vapour pressure of solvent in solution

 $p_{\phi} = N$ apour pressure of pose solvent.

 $z_i = Mole fraction of solvent.$

$$z_k + z_k = 1$$

Where $x_{ij} =$ mole fraction of solute

Then

$$z_t = 1 \cdot z_t \perp (2)$$

Put the value of x_k from eq.(2) in eq.(1)

$$P = P^*(1-x_0)$$

 $P = P^{\alpha} \cdot P^{\alpha} x_{\alpha}$

 $P^k \circ P = P^k x_k$

$$\Delta P = P^{\bullet} \pi_{0} (3)$$

Where AP = Lowering of Vapour presons

 $\Delta P \propto z_0$

Thus, Repults law our also be stated at

Emerged of supers has

to Mathematically AP a Te

From eq (3) we have

Where $\frac{\Delta P}{P^2}$ = Ratetive lowering of vapour pressure

Relative lowering of exposer pressure of a solution is equal to the mole fraction of man Facts about relative lowering of Vapour Pressure

- It is independent of the temperature
- it depends upon the conc. of solute
- It is some when equippolar amounts of different solutes are dissolved in the more amount of the same solvent,

Result's Law and Ideality of solutions

A solution, which obeys Raouit's law, is called an ideal solution. Thus, Raouit's law as help to distinguish the ideal and non-ideal behaviour of solutions,

The capour pressure of uniter at $30^{\circ}\mathrm{C}$ is 28.4 tors. Calculate the suppour pressure of a solution containing 70 g of came sugar $(C_{12}H_{22}O_{11})$ in 1000 g of water at the same temperature. Also calculate the lowering of suppour pressure.

Mass of Cane Sugar = 70 g

Molar Mass of Cane sugar (C $_{tz}$ H $_{zz}$ O $_{11}) \quad = 342 \ g$

Moles of case sugar = $n_2 = \frac{70}{342} = 0.20$ moles

Mass of water = 1000 g

Motor Mass of water = 18 g

Motes of water = $n_1 = \frac{1000}{19} = 55.5$ moles 18

Thus Mole fraction of came sugar $\Rightarrow x_0 = \frac{0.2}{0.20 + 55.5} = 0.0036$

Mole fraction of water = $x_1 = \frac{55.5}{0.20 + 55.5} = 0.9964$ 55.5

Mole fraction of water $= x_1 = 0.9964$

Charleton Part-i

512 Vapour pressure of pure water ≈ 20.4 tor Vapour pressure of solution = P = 2 According to Repull's law $P = P^* x_1 = 28.4 \times 0.9964 = 28.29 torr$ So lowering of vapour pressure = 28.4 - 28.29 = 0.11 tour

January Lane for two Volume Components

In a solution of two solutile components A and B the sepons presents of each onent above the solution is proportional to the mole fraction in the solution.

$$P_A = P_A^a x_A \qquad P_B = P_B^a x_B$$

Where $P_A = Vapour$ pressure of A above solution

Pa = Vapour pressure of B above solution

P = Vapour pressure of pure A

P° = Vapour pressure of pure B

 $x_A = Mole fraction of A$

 $x_n = Mole fraction of B$

According to Dalton's law, total pressure of the colution is equal to the sum individual pressures of A and B

$$\theta_i e_i = P_i = P_A + P_B$$

$$or \qquad P_i = P_i^a x_a + P_i^a x_a$$

 $\kappa_A + \kappa_B = 1$ or $\kappa_A = 1$ κ_B Since

Hence
$$P_i = P_A^a (1 - x_B) + P_B^b x_B$$

$$P_1 = P_{A}^a - P_{A}^b x_b + P_{B}^b x_b$$

$$P_k = (P_{ij}^a P_A^b)x_i + P_A^b$$

 $P_k = (P_{ij}^a P_A^b)x_i + P_A^b$
 $P_A^b = P_A^b$, and intercept = F

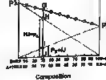
This is an equation of straight line with slope $P^a_{\ A}=P^a_{\ A}$ and intercept $=P^a_{\ A}$

Only ideal solutions will give straight lines.

Graph shows following things

- Dotted lines represent the vapour pressure of each component in the solution, which increases with mercase in mole fraction.
- * The vapour pressure of solution is given by

the straight line joining $P^{a}_{\ A}$ and $P^{b}_{\ B}$





College Chambios Paris!

5.1

- Vapour pressure of solution is never greater than P⁶_A or less than P⁶_— Let it remains intermediate between $P^a_{\ A}$ and $P^a_{\ B}$
- Vapour pressure of each component in the solution. is always less than its vapour pressure in pure form.
- Total Pressure of sociation is equal to the sum of individual pressures of components A and B in the solution.

e.g. consider the point G in the fig. At this point vapour pressure of A is $P_A[H]$ and that of B is Ps (U). Thus, total vapour pressure of solution at G will be $P_{i} = P_{A} + P_{B} = HU + U$

THE ALL AND NON THE ALL SOLUTIONS

then! Salutions

- (1) A solution, which obeys Rapult's law, is called an ideal solution,
- (2) During formation of an ideal solution, heat is not evolved or absorbed i.e., enthally change is zero. $\Delta H = 0$
- (3) Total volume of solution is equal to the sum of volume of components of solution.
- (4) Forces of attraction among molecules of components remain same in the solution just as they were in pure form

Exemples

Bertzene - Toksene, Benzene - Ether, Chlorobenzene-Bromobenzene, Ethyl iodide Ethyl bromide etc.

Southbut Seduces

- (1) A solution which does not obey Result's law is called a non-ideal solution.
- (2) During formation of a non-ideal solution, heat may evolve or absorb ..e., enthalty change is not zero. $\Delta H \neq 0$
- (3) Total volume of solution is not equal to the sum of volume of components of solution.
- (4) Forces of attraction among molecules of components do not remain seme in the solution just as they were in pure form.

Examples.

Ethanol-Water, HCI-Water etc.

Difference between Ideal and Non-Ideal colutions Ideal Solutions

976

A solution, which obeys Recult's law la called an Ideal solution

During formation of an ideal solution, heat is not evolved or absorbed up. enthalpy change is zero (AH=0)

Total volume of solution is equal to the sum of volume of components of solution.

Forces of attraction among molecules of components do not remain same in the solution just as they were in pure form.

Benzene Toluene. Benzene - Ether Ethyl lodide - Ethyl bromide etc.

Non-alcal Solutions A solution which does not obey Reput's law is called a non-ideal solution.

During formation of a non-ideal solution,

heat may evolve or absorb. Le., enthalpy change is not zero (AH=0)

Total volume of solution is not equal to the sum of volume of components of solution.

Forces of attraction among molecules of components do not remain same in the solution just as they were in pure form.

Examples:

Ethanci-Water HCI-Water etc.

VAPOUR PRESSURE OF LIQUID-LIQUID SOLUTIONS

Exercise QB. (a)

Explain fractional distillation, Justify the two curves when composition is picted against bolling point of solutions.

ideal Salution

A solution which obeys Recult's law is called an ideal solution.

Proctional Distillation

The superation of liquids from a liquid mixture on the bests of their boiling point is talked Prantished distillation.

n (thum) Distillation of Ideal Schillar

Consider a solution of two votable components A and B. Let A is more votable than B. hence A has lower boiling point than B

Let solutions of various compositions are made and composition of estution and its specing is plotted on X-axis against their boiling

herefure on Y-axis Thus, two curves are obtained as shown in the fig.

- Lower curve gives the composition of solution at different temperature, while
- * Upper curve gives the composition of vapours in Equilibrium with solution at different tempera









Two curves are obtained because vapouss and solution have different composition a different temperature.

When such liquid mixture is heated to point G. then

- Composition of solution is given by C which have more B and less A, while
- Composition of vapours is given by D which have more A and less B

Since A ta more volatile, therefore at each temperature variours have more % of A than B. The liquid mixture left behind is collect residue and the superer abiolysed ers cally distribute.

The distillate obtained is cooled and again distilled. The re-distilled vapours will belone notes in A then before and if this process is repeated many times then various will be on notes in A and residue will go on notes in B. Hence both A and B can be separated. Aut peer over while B will be left behind. Such a mixture is called zeotropic mixture.

Ligade which ducies with change in composition are called scotrapte missures.

Zeotropic mixture can be easily separated into their components.

Example.

Methyl alcohol -- water solution can be separated by distillation.

Extrelas Q7

What are non-ideal solutions? Discuss their types and give three example of each Exercise Q S(b)

The solutions showing positive and negative deviations cannot be fractionally distilled at the specific compositions. Explain it,

What are assorropic mixtures? Explain them with the help of graphs?

See Adeal Schuttones (Azentroque Michaels)

A retuitor, which does not about Requir's law, is sailed a non-ideal solution.

They show deviations due to difference in their molecular structure i.e. size, shape and intermolecular forces.

Appointante Ministree

The liquid mixtures, which had at openions temperature and distil over stakes change in composition like a pure compound, at any temperature are called applicable mixtures.

These are not true chemical compounds. Because their composition can be changed by changing conditions such as pressure. While composition of pure chamical composition to pure chamical composition. termains constant at different temperature and pressure.

College Chambatric Part-I

421 (metronal distillation of New-Ideal Sedations

Non-ideal solutions show either maximum or minimum in their vepour pressure curve. Hence, they connot be separated completely by fractional digilitation.

Non-ideal solutions show two types of deviations

- Positive deviations
- Negative deviations

Vapour pressure-composition diagram for both types is shows in the Ba.



Poeliine Deplatione (Minimum Bollino Agarigose)

Liquid solutions which have maximum in their vapour pressure curve shows position

Consider a solution of two components A and B. in which A is more volatile than B. In lig, at point C. vapour pressure of solution is maximum. Thus, at this point boiling point will be minimum. Hence, on distilling, that this mixture of constant composition is distilled over then the temperature will rise and other component will pass over

in such solutions, complete separation of components is not possible. Only a pure component and a minimum boiling azeotropic mixture is obtained.

The exectropic minture boiling at minimum temperature than any of its component is called musimum boiling assotrops Example

Ethaniol - water forms assotropic mixture with ethanol 95.6% it both at 78.1 °C This imperature is less than the boiling point of both ethenol (78.5°C) and water 100°C)

Henning Deviations (Maximum Sailine Assetrops)

Liquid solutions, which have minimum in their separa pressure cured, should negative riottona from Reputt's loss

Consider a solution of two components A and B, in which A is more vowille than B.

In fig. at point D the vapour pressure is minimum, hence, belling point is maximum. As his Point liquid mixture distills over without change in composition. It is called maximum bolling assetrope.

Thus, two components cannot be separated. Only one component and a constant folling executopic mixture is obtained

The temotrope halling at maximum temperature then any of its compound is called Maximum boiling assotrop

Emple

tk] - Water postern

HCL forms azeofropic mixture with water, which bolls at 110°C. It contains 20.24 % HCL.









Security of Solution

The aphytion of

has donated solution.

ch more solute can be disvalped at à given temperature is colei

Sofulation

It is the concentration of solute in solution when it is in signification with the edit retance in its enteroise solution at a given temperature.

Solubility is expressed as number of g, of golute in 100g of solvent.

Generally solubility increases with increase in temperature

At O°C, solubility of NaCl in H₂O is 37:5 g / 100 g of H₂O

At 0°C, solubility of CuSO, in H_gO is $14.3~\mathrm{g}$ / $100~\mathrm{g}$ of H_gO

At 100°C solubility of CuSO₄ in H₂O is 75.4 g / 100 g of H₂O

When a solute is dissolved in a solvent, its molecules or lone break away and disense in the solvent. This is called <u>dissolution</u>.

If solution becomes substated with solute, then lone or motecules may re-unity to form precipitate This is called re-crystalization or precipitation.

In a solurated solution, a dynamic equilibrium is established between solid solute and dimolved solute. At this stage rate of dissolution becomes equal to the rate of crystallization. At this temperature, maximum amount of solute dissolved in solvent is called solubility.

Determination of whilatar

- Plast, a saturated solution of solute is prepared in a given solvent.
- Solution is filtered
- A known volume of this solution is evaporated until solid regidue is left behind
- From the mass of residue and volume of solution evaporated, solubility can be calculated

Solubility Correct

A graphical representation between temperature and solubility of a substance is wild solubility ourse.

These may be of two types

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(1) Continuous Schubtly Curpes

These are smooth curous and do not show a

These may be of two types

(i) Increasing Solubility Curve

(ii) Decreasing Solubility Ourses

(Il-Increasing Soublility Curve

Formany substances increase in temperature

increases the solubility.

Thus increasing solubility curve is obtained.

Example

Solubidity curves of KCIO₁ K₁C₂O₂ Pb(NO₃)₂ and GaCl₄ Solubility curves of NaCl shows only alight increase from 0°C to 100°C

M.Decreasing Solubility Curbes

For scane sails, increase in temperature decreases the solubility of sait.

Thus decreasing solubility curve a produced,

Example

Solubility of Ce₂(SO₄)₃ decreases with increase in T upto 40°C. After 40°C it becomes Imost constant.

Discontinuous Salubility Curves

These are not smooth and should saiden breaks

due so eudden ahanges in solubilities

Pierrole .

Solubility purver of Ne₂SO₄, 10H₁O, CeO₆, 6H₂O etc.

in these cases, actually a new phase appears at the brak point

Thus, next curve is the solubility curve of this new phase

Mance discontinuous solubility curves are actually dombinations of two or more continuous solubility curves

actional Crystallization

The esparation of solid substances from a solution, one by one, on society to called rectional organollisation.

Prominence: It is a technique for separation of impunities from chemical products.

Solubilities of substances depend upon temperature

4.2. solubility of KNO, rapidly changes with temperatrus but solubility of KCI, KBr



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1014

Atanine etc changes gradually.

Thus one substance may precipitate earlier by cooling, leaving behind other in the solution. This is the basis of Fractional Crystallization.

Mached impure solute is dissolved in hot solvent. In this solvent, solute must be less solute than impurities,

As the hol solution is cooled, pure solute crystallizes out first leaving impurities in the solution. Thus crude product can be purified.

Exercise O10, (a)

What are the colligative properties? Why are the called so?

COLLIGATIVE PROPERTIES OF DIFFIT SOLUTIONS

The properties, which depend upon number of particles and not on their nature, or olled colligation properties

Examples

Lowering of Vapous pressure, Elevation in boiling point, Depression in freezing point, Osmotic pressure

Lydanation: Why some properties are colligative?

Pure water has definite vapour pressure at a given temperature

Suppose 6g of area, 18g of glucose and 34.2g of sucrose are dissolved in 1 kg of Hi^O. Each of this is 0.1 motal solution and contain equal number of solute particles, i.e. 1/10 ft of Avogadro s particles i.e 6.02 x 10 22

In all these cases, lowering of vapour pressure is same. The boding point of solution is rained by 0.652%C, and freezing point of solution is depressed by 0.186%C than pure H_0O

Thus towering of vapour pressure, Elevation in boiling point and Depression in Freeing point are some in all cases due to some number of particles. Hence, these properties of called colligative properties.

Exercise Q10 (ii): What is the physical significance of K_b and K_f values of solvents?

Molal Boiling Point Constant or EhalDoscopic Constant

It is the elevation to holling point, which is produced, when I make of epide ed in I kg of solvent.

It is denoted by $k_{\rm h}$

if I make of solute e.g. uses (50 g), or sucrose (342 g) or glucose (180g) is dispolved in H₆O. The boiling point elevation of water is 0.52 °C. Hence is, for water is 0.52°C

Significance: It tells about how much elevation of boiling point will occur for a period motality of schulon, e.g. for water the elevation in b.p. would be 0.52°C/maid solution.

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Hold Freezing Point Constant or Cryoscopic Constant

to the depression to freezing point which is produced, when I make of actual is

) is denoted by k_i .

If mote of solute e.g. unsa (60 g) or suggest (342g, or glucose (180g) is discoved in R_0O . The depression in freezing point of water is 1.86°C. Hence, R_0 for water is 1.86°C.

Significance: It tells about how much depression of freezing point will occur for a particular molality of solution. e.g. for water the depression in f.p. would be 1.86°C/molal solution.

Table (9.3) K, and K, values for some salvents							
Solven .	BiPales	· 秘密學 mir-	· 我见 ///3	P. (No. 2 M. Pem)			
H ₂ O	100	0.52	a	85			
Elher	34.4	2.16	-116.3	. 79			
Acetic actd	116	3.07	17	3.90			
Ethanoi	79	175	14.5	199			

Conditions For Normal Colligative Properties

- Solution must be dilute
- Solute must be non-volatile
- Solute must be non-electrolyte

Sudy of colligative properties has given a method of molecular mass determination. This and also developed the solution theory.

How do your explain that the lowering of capour pressure is a calligative property? How do you measure the molar mass of a non-volatile, non-electrolyte solute in a palent. 917 polotila solvent?



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For eving of Vigorov pressure When a non-colonie, non-electrolyte solute to dissolved in a volatile solum, a sour presents to losswed. This is called lossering of capour presents.

In pure solvent, all the surface of solvent is covered by solvent molecules.

But when a non-volatile, non-electrolyte solute is paded to it to form solution. Solutions is no coupled by solute particles. Harves escaping tendency of solvent is decreased thus its vapour pressure is covered.





Relationship to a foregroup of renount pressure multipuly traction of solute

tion: Determination of Mal. More of Solute).

According to Recuits law

Enlettee temesting of persons pressure is equal to mote fraction of solute.

$$\lim_{n\to\infty}\frac{\Delta^n}{p^n} *_{X_n^n} \underline{\hspace{1cm}} (1)$$

△P = Lowering of vapour pressure

P* = Vapour pressure of pure solvent

 $n_{\rm d}$ = Male Fraction of solute

$$\chi_2 = \frac{n_2}{n_1 + n_2}$$

Where $n_{\rm s}$ and $n_{\rm f}$ are the number of moles of solute and solvent respectively For dilute solutions, ne is very small, therefore it can be neglected in the denorite

Since $n_1 = \frac{W_1}{M_1}$ and $n_2 = \frac{W_2}{M_2}$, therefore

$$h_2 = \frac{W_2}{M_*}$$
 , therefore

$$\underline{w}_2 = \frac{n_2}{n_1} = \frac{W_2 M_1}{W_1 M_2}$$

$$W_g = Mass of solute (g)$$

 $W_i = Mass of solvent(g)$

 $M_k = Mol.$ Mass of solvent $\{g\}$

 $M_e = Mol$, Mass of solute (g)

$$\frac{\Delta P}{P^0} = \frac{W_2\,M_1}{W_1\,M_2}$$

or
$$M_2 = \frac{p^0}{\Delta P} \times \frac{W_2 M_1}{W_1}$$

Using this equation moterular mass of solute "M" can be determined.

Example 10

parties for a suppose pressure of 122 tors at 12°C. When 20 g of a con-admits were discolard in 200 g of bensens, a suppose pressure of 120 core was observed. Calculate the molecular mass of the volute. The molecular mass of bensents is 78.1 giand.

 $-w_1 = 20 g$ Meas of solute "X"

Meas of solvant banzene = w_s = 300 g

Moler Meas of benzene = M_s = 78.1 g/mol

Vapour pressure of pure after = P⁴ = 122 tor

Vapour pressure of pure other $= Y^* = 122$ for Vapour pressure of solution = P = 120 for Lowering of vapour pressure $= AP = P^0 - P = 122 - 120 = 2$ for Moter Mass of 'X' $= M_X$ = 2

Molar Misso of solute is given by

$$M_2 = \frac{P^0}{\Delta P} \times \frac{w_2 M_1}{w_{1.0}}$$

$$M_2 = \frac{\Delta P}{\Delta P} \times \frac{\omega_{1.6}}{\omega_{1.6}}$$
 $M_2 = \frac{122}{2} \times \frac{20 \times 78.1}{300} = \frac{317.6 \text{ g/mol}}{2}$



Entetes Q14

Out graphical explanation for elevation of boiling point of a solution. Describe one method to describe explanation for elevation of a solution determine the boiling point elevation of a solution

bouling of Boiling Point The difference between the boiling points of the solution containing non-volatile non-incipality and the pure solvent is known as boiling point alsoution of the solution, and the pure solvent is known as boiling point alsoution of the solution.

It is denoted by ΔT_{ϕ}



hat Charles: Part-I

52.4

A Equid both at a temperature at which his cape tensor equal to the automol pressure.

Let AB is the vepour pressure-temperature curve of pure solvent. This graph is roade by measuring the vapour pressure of pure solvent at different temperatures.

When a non-volatile solute is added to the solvent, the vapour pressure of the resulting solution is lowered, therefore, vapour pressure-temperature curve of solution, CD lies below that of pure

Let external presoure is P*.



It is clear from the graph that solvent attains this pressure at T_{\parallel} and starts below However, notation sitiates this pressure at T_2 , which is higher than T_3 . Thus, solution toles higher temperature than pure solvent.

Such that $T_A - T_A = AT_A =$ Elevation in botting point

Generally, higher the conc. of solute, greater will be the lowering of vapour present Hence higher will be the elevation in boiling point.

Thus elevation in holiting point is directly proportional to molality of solution \boldsymbol{x} $\Delta T_b = \mu_b \times m$

Where

ATh - Elevation in boiling point

k_k = Exultoscopic constant

m = motality of solution

According to eq (1), 6 g of uses or 18 g of glucose in 500 g of water reach is 0.2 not aduation) have seens elevation of boiling points. I.e. 0.1°C which is 1/5 th of 0.52°C 5c life is a collection of the co is a colligative property

Since

$$\begin{aligned} & \text{Motelley} = \frac{\text{Mass of solute}}{\text{Molecular Mass of solute}} & \frac{1}{\text{Mass of solvent in Kg}} \\ & \text{fo} = \frac{W_3}{M_2} \times \frac{1}{W/1000} \\ & \text{or} & \text{m} = \frac{W_3}{M_2} \times \frac{1000}{W_1} & \text{(2)G} \\ & \text{Where} & W_8 = \text{Mass of solute (g)} \\ & W_1 = \text{Mass of solute (g)} \\ & W_8 = \text{Moter Mass of solute (g)} \\ & M_8 = \text{Moter Mass of solute (g)} \end{aligned}$$

Put eq (2) in (1)
$$\Delta T_b = \kappa_b \times \frac{W_2}{M_2} \times \frac{1000}{W_1}$$

or
$$M_2 = \frac{k_b}{\Delta T_b} \times \frac{W_2 \times 1000}{W_1}$$

Using this eq. Molar Mass of solute ($M_{\rm p}$) can be calculated.

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Measurement of thermion of Builting Point (Landsberger's Method)

in this method, solution is heated by vapours of bailing solvent

Apperatus

It consists of

An inner botiling graduated tube with a hole in the side

A boiling flask, which sends vapours of solvent to the inner tube through a rose head.

An outer tabe, which receive vapours of hot solvent through the hole of inner tube.

A Beclaneum thermometer that can read upto 0.01 K.

A small amount of solvent is placed in the inner bosing tue. It is boiled by the vepours of scheet coming from toding fiels. These vepours boil the selvent in the tube by their latent hast of sondemention. When solvent boils its boiling point is noted.

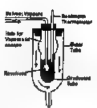
Now supply of vapours is out off. A weighed quentity of solute is added to the solvent in the inner table. The boiling point of solution is also noted similarly. The difference of boiling point of solution and solvent gives the slevation in boiling point.

After noting boiling point, volume of solution is also noted to calculate the mass of loivent (sv.

Molar Mass of solute can be determined by the formula

$$M_2 = \frac{M_b}{6 T_b} \times \frac{W_2 \times 1000}{W_3}$$

hample II ing point of moter is 99.785°C. To a sample of 600 g of wa Wing molecular mass 88 gimel, to form a solution. Calcular





Charles from

Mass of solute:
$$\infty$$
 w_p ≈ 24 g

Mass of solvent water: ∞ w_k ≈ 600 g

Mobil 8.P construct of water= k_p ≈ 0.52 °C

Mobil Mass of solute= M_p ≈ 58 g/mot

8.P of pass water: ∞ T = 99.725 °C

Execution in 8.P \approx AT_p \approx ?

Elevation in boiling point is given by

$$\Delta T_{b} = i_{b} \times \frac{m_{Z}}{M_{Z}} \times \frac{1000}{m_{Z}}$$

$$\Delta T_{b} = 0.52 \times \frac{24}{58} \times \frac{1000}{600} = 0.368 ^{\circ}\text{C}$$
Hence B.P. of solution = $T^{\circ} + \Delta T_{b}$

$$= 99.725 + 0.358 = [100.080 ^{\circ}\text{C}]$$



Emercine Q15:

Freezing points of solutions are depressed when non-valuable solutes are present in with solvents, shadify it. Plot a graph to elaborate your answer Also, give one method to real the depression of freezing point of a solution.

Depression of Feerzing Point

The difference in the freezing point of equation containing non-solable, and electrosyste solute and pure actions is called depression in freezing point.

It is denoted by ΔT_f

Freezing point of a liquid to defined so the T at which told and liquid phases have some capour pressures.

At freezing point both liquid and solid phases are in

Consider the vapour pressure—temperature curves of pure solid solvent BC (sublimation curve), pure liquid solvent 'AB' and solution 'ED' as shown in the fle.

Vapour pressure curve of solid solvent is more steen

showing that the change in vapour cressure is more round with termetrature.



Carrier Charles Con-1

Stage vapour pressure of solution curve is lower than that of pure solvent, therefore, in pour pressures immigrations curve is below that of pure solvent.

At Sensing point of solvent both solid and liquid phases are in equilibrium and have size exposur posseure. It occurs at point B, to it is the integing point of pure solvent.

At freezing point of solution both solid tolerat and boud solution are in confibring and the same vapour manage. It occurs at point E. So it is the freezing point of solution.

Since $T_1 > T_2$, therefore solution freezes at lower temperature that that of ours solvent.

$$\Delta T_f = T_S \cdot T_g$$

The deposition in freezing point is directly proportional to moisity of solution i.e. AT, C M

$$\begin{array}{lll} \Delta T_t = |t_d \times m| & \text{in} \\ \text{Where} \\ \Delta T_t = \text{Depression is freezing point} \\ k_t = \text{Cryoscopic constant} \\ m = \text{Moleity of solution} \end{array}$$

Since Mobility =
$$\frac{\text{Mass of solute}}{\text{Mobility Mass of solute}} \times \frac{1}{\text{Mass of solvent in Kg}} \text{ otherwise}$$

$$m = \frac{W_2}{M_Z} \times \frac{1}{W_3} \frac{1}{1000}$$
or
$$m = \frac{W_2}{M_Z} \times \frac{1}{W_3} \frac{1000}{W_2} = \frac{1000}{W_2} \frac{1000}{W_3}$$

Using this equation Molar Mass of solute ($M_{\mbox{\scriptsize g}}$) can be calculated.



Collage Chambers, Part-I

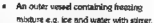
Measurement of Depression in Freezing Point (Backmann's method) Apparatus

It consists of

· A freezing tube with a side arm fitted with a Becommen thermometer. It has stirrer and contains

solvent or solution.

An outer tube surrounding the inner tube Space between outer and inner tube behave as air jacket for slower and uniform rate of cooling.





About 20-25 g of solvent is taken in the freezing tube. The bulb of thermoresers dipped in the solvent. First, an approximate freezing point of solvent is determined by directly cooling the tube in freezing mixture

Freezing tube is then taken out. After re-melting solvent, it is then piaced in air jackrise cooled slowly and accurate freezing point of solvent is noted.

Now about 0.2 g to 0.3 g of solute is added to the solvent. The freezing point of solution is determined by stirring the solution

The difference of freezing points of solution and solvent gives the depression in [seem]

Moiar Mass of solute can be determined by the formula

$$M_2 = \frac{k_f}{\Delta T} \times \frac{W_2 \times 1000}{W_1}$$

Example 12

The freezing point of sure compher is 178.4°C. Find the freezing point of a science containing 2 g of a non-velocitic compound, having melecular mass 160, in 40 f 4 compher. The moial freezing point constant of compher is 37.7 °C kg /moi.

Solution

Mass of solute compound = w_e = 2 g Mass of solvent camphor $= w_1 = 40 g$ Motal F.P. constant of campho: = k_t = 37.7 ^aC. kg/mol Molecular Mass of solute = M₈ = 140 g/mol

Chemistre Parts

Freezing point of pure camphor = T = 178.4 °C Freezing point depression $\Rightarrow \Delta T_f = ?$ Freezing point of solution = ?

Depression in F.P. is given by

$$\begin{split} &\Delta T_f = |_{k_1} \times \frac{w_2}{M_2} \times \frac{1000}{w_1} \\ &\Delta T_r = 37.7 \times \frac{2}{140} \times \frac{1000}{40} = 13.46 ^{\circ} C \\ &\text{Hence F.P. of solution} = T^{\circ} - \Delta T_r \\ &= 178.4 - 13.46 = \underline{164.94 ^{\circ} C} \end{split}$$

Applications of B.P. Electrica and L.P. Depression

Imperation of antifreeze

in winter water in radiators of automobile may freeze. To protect this a solute ethylene styrol is added to it. It is completely miscible with H₂O. It is also almost non-volatile. Thus it depresses the freezing point of H_2O Hence, it prevents the freezing of water in winter

In summer it protects the radiator from boiling over

P<u>tenarotion of Freesias mixture</u>

When NaCl or KNOs is added to Higo, its imasing point is lowered. This mixture is used a frezing muciure in an los-cream machine.

Estroire Q16:

Discuss the energetics of solution. Justify the heats of solutions as exothermic and indoshermic properties.

(MRGLHES OF SOLDHON)

The amount of heat absorbed or evolved when a substance to described in a softent to Is my a solution to called heat of solution.

The mithalpy of a solution of a substance is defined as the first changed when one ide of a substance is described in a specified number of moise of solvent at a given in the substance is dissolved in a specified number of moise of solvent at a given

It is denoted by $\Delta H_{\rm sole}$. It is the heat of solution at infinite disution Sample

When 1 more of NeCl (58.5g) is dissolved in 10 moter of H_sO (180g), 2,008 kJ energy is other bedroad

ΔH ≠+ 2,008 Ke/mol $N_B Cl_{(i)} + 10H_B O \longrightarrow N_B Cl(10H_B O)$



Exploration

(i) When a solute is dissolved in solvent, following things occur Molecules of solvents move esvay from each other. This process requires energy to overcome the cohesive intermolecular forces, hance it is endothermic

(ii) Solute molecules separate from each other. This process also requires energy hence it is also endothermic.

(iii) Solute and solvent particles develop forces for each other and energy is released. It is an exothermic process.

The exothermic or endothermic nature of process depends upon the strength of two types of forces.

The process of dissolution occurs with either absorption or release of energy. This is due to breaking and remaking of intermolecular forces between solute and solvent

When KNO₄ is dissolved in water, Temperature is decreased showing endotherize

When LICI is dissolved in Pi₂O, Temperature is increased. Thus it is exothermic process. Mathematically

 $\Delta H_{\rm sole} = H_{\rm columbs} - H_{\rm so}$

Where $\Delta H_{\rm soft}$ = Enthalpy of solution

H_{relegion} = Heat content of solution after formation

H_____ = Heat content of components before rubing

Both H_{sequents} and H_{states} can not be calculated

Only AH_{eb} can be calculated

ΔH_{set} is -ve when H_{sets} < H_{sets}

AHees b +ve when Harry > Harry

hillian entern of ente

When an lonic compound is dissolved in solvent. Some forces develop between long of and golvent. This concess releases to the solvent in solvent. solid and solvent. This process releases charge, This energy, realistsed ultimates breaks had been colin and it discourse. lattice of lonic solid and it dissolves.

page de

Table: Hears of solutional

4.98

26.0

17.8

21.4

NaC

NHLNO,

KCL

ĸ

Dissolution of KI in H_sO takes place in two steps

in first step lattice of KI is broken down due to solute . solvent interaction. This process requires energy hence it is enciothermic. The energy required to break lattice is called lattice energy.

The process is

lattice energy of route solids is always meater than molecular solids. Amount of the more



* + 11	4.00
to the same	01 8an
H: Li ^b No+	~1075
Ţ,	-499
No*	-390
Ag*	464
K* .	-305
Mg ²	
Crit	1691 1562
NH,+	-281
P-	-457
<u>a</u> -	-384
NH ₄ °	-851

In second step, K* and I* ions produced are surrounded by dipoles of water. This process is called hydration or solvation. During this process, energy is misseed, it is called heat of hydration, e.g.,

Thus, overall process is

$$KI + xH_{A}O \longrightarrow K^{*}_{pap} + \Gamma_{pag}$$

Different ions have different heats of hydration

Sea of lank size on Hydration Energy

Generally greater the ionic radii of menoposities ions, smaller is the heat of operation

Divalent and Trivalent ions have greater heat of hydration due to large charge densities.

Anions also show similar trend.

Infinite Offution

On adding more H_eO to NaCl solution, <u>further heat chance</u> may occurs. This heat change depends upon the amount of water used for distion. The heat of distion stows decreases and finally no further heat change occurs. This occurs, when there are \$20 - 1000. The is called introduced to the control of the cont

Thus heat of solution at infinite dilution will be

Δ H_{mb} = +4.90 kl/mol Napal++Cleat NaClus + HO - NaClus



OVERATION AND DATERUASIS

DVDRAHON

The process in which water molecules surround and interest with the solute parity, both breaking my O-H band is called hydration

Limitensiine

Water is an excellent solvent, it dissolves various lonic substances because of hydrates of ions. A hydrated ion is a cluster of ions and one or more water molecules.

in solution, each ion is surrounded by indefinite no. of water molecules.

However when water solution of a salt is evaporated, the sait crystallizes out with definite no of water molecules. These water molecules are called water of crystallization and are present in crystal lattice of sait. During this process no bonds are broken. However new bonds are formed

in Janus substances, negative ions are tileger and positive ligne are smaller in size. The manuscript is size in a size of the charge control with a positive ions have low charge density (charge per unit volume), while positive ions that high charge density.

Generally. Ions with smaller size and high charge dinsity — e. +toe ions , have not attraction for polar water molecules than \sim 0 tons. Thus, in CuSO₄, SH₂O, but of five wife molecule, four are attached with smaller Cu⁸⁺ ion and one is attached with SO₄** ion

Distance

The expetalline substance, containing abenically combined $\mathcal{H}_2\mathcal{O}$ is, define proportione is called hydrate

Examples

CuSO, 5 H,O Na,SO, 10H,O

In CuSO₄ 5H₂O , four H₂O are attached with Cu¹⁺ and one with SO₄ **

Company of High Street

Hydrates are prepared by gyaponating their nameous solutions.

Hydrates are not limited to salts. Acids, bases and elements also form hydrates.

A CONTRACTOR OF SIG

The senter molecules, which we attached suith compound when espetalized frost of setter, are called noter of espetalization.

Entropie

(COOH)₈ 2H₄O (Oxalic acid) B₈Cl₄ 2H₄O

MgCla 6HaO

MgSO₄ 7H₂O (Epsom salt)

NaCO, 10 H₂O

CeSO₄ 2H_gO (Gypaum)

Na₂B₄O₇ 10H₂O (Borney)

Callege Chartisters, Purch

SPROUSIS

The process in which mater reacts shemically with the edded solute to calle

<u>Reviewation</u>

A said produces cations and anions or dissolving in water. These long may react with water to produce scidic or basic solution

Cation hydrolysis produces addic solution

M*+ H-OH - MOH + H*raddic solution)

Anion hydrolysis produces basic solution

A" + H-OH - + OH" (basic solution)

Generally sails may be dassified into four types.

17 THE PARTY AND DESCRIPTION AND DESCRIPTION OF PARTY AND DESCRIPTION O

e.g. CuSO₄ .lt lonteer as

CuSO₄ Cult + SO₄*

Thus eq. solution of CuSO₄ is addic.

Other Examples

A(C)₅ + 3H₈O A((OH)₆ + 3H* + 3C|* NH₄Cl + H₈O NH₄OH + H* + Cl*

Thus, these hydrolysis reactions produce weak bases and strong acids. Both Cl. and are week conjugate bases of strong acids HCl and H₂SO₆ respectively. Thus they are for hydolysed by water. The H° are free in solution, therefore, their solutions are acidic in Patture. The K_a values are of HCI and H₂SO₄ are very high as compared to K_a values of Al(DH)₈, NH₄OH and Cu(OH)₈.

2. Sales Of Strong Boso With Week Acid.

P.B. CH₈COONa It ionlass as

CH_aCOONs = CH_aCOO*+ Na*

CH₈CDO hydrolyse in water to produce basic solution

CH,COO+ H-OH ____ CH,COOH + OH

Thus eq. solution of CH₂COONs is basic.



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Other Exemples

NewPOat, NewAnDards

These also give basic solution due to formation of Na* and OH " and weak acids Hapo, and H_eAsO_e respectively, which are not ionized.

CONTRACTOR OF THE PARTY ACT

By adding such sait in water, resulting solution may be acidic, basic or neutral depending upon the pK, and pK, values of said and base produced.

CH_COONH_ NH_CN es.

- San Of Server and Ultra Procession

a.g. NuClionius as

NeCl 🚤 Not + Cl *

Both No.* and CT are not hydrolyzed by H₂O, therefore, it's solution will be neutral. Thus sails of strong acids and strong bases are not hydrolyer-1 by water.

Other Emerales

Na₂SO₂ , RBr, RCI etc.

Thus, sails of strong unid and strong bases are not hydrolyzed by water

Deference between Hydration and Hydrolysis

DADRATION The present in which water molecules surround and interact with the solute purchin without breaking any bond is alled hydration

Largely physical interactions occur en solum and water molecules.

Old bonds are not broken but new bonds may be formed.

This process does not change the pH of

INDROLYSIS

The process in which water reads chemically with the added solute is called hydrolysis

Chemical reaction occurs between source particles and water molecules
Old bonds are broken and new book

This process may change the phi of solution.

Charles Park

300

(d) 6

OBJECTIVE AND SOLORED SYSWER OF ESTE AS TALIN (8) Multiple Choice Questions

Molectly of pure weter to

(b) 18 (a) 1

kl 55.5

(6) A (1) 5

(6) 5

(6) 5

(6) 5

(6) 5

(7) 5

(8) 6 Sound 2019 (Rosenjahad Sound 2010) (R.G. Nime Sound 2011) (Solate Sound (a) 1/5 (b) 5.1 (c) 1/51

perf board, 2012) (Calemania board, 2008, 2019)

(a) A solution of plucose is 10%. The volu o to which I greate of it to die of will be (a) 1 dm2 (b) 1.8 dm⁸ (c) 200 cm²

(d) 900 cm³ p.G. (Dan Board, 3712) (Sergodia Seard, 2012)

(v) An aqueous solution of methanol in unior has repose presented fall equal to that of water (b) equal to that of methanol (c) more than that of water (d) less than that of water

apur Board, 2010) (Sorgadia Board, 2013)

An attotropic mixture of two liquids both at a le them when (Facilities Series, 2027)

(a) It is saturated

(b) it shows positive deviation from Rapult's less

(c) It shows negative deviation from Racult's lew

(d) It is metastabia

(40) In anestropic maxture showing positive deviation from Result's law, the value the minture lo

(a) Slighlify more than the total volume of the coreponents

(b) Slightly less than the total volume of the components

(c) Equal to the volume of the components

(d) None of these

(vi) Which of the following solutions has the highest beiling point?

(a) 5.85 % solution of NaCl (b) 18 % solution of glucose

(c) 6 % solution of unan (d) all have the same bolling point

with Two county of NaCl and KCl are properly apparately by dissolving wants and the solutions of NaCl and KCl are properly apparately by dissolving wants as the solute in water. Which of the following shatements is true for these soluted by solution.

(a) KCI solution will have higher boiling point than NeCl solution

(b) Both the solutions have different boiling point.

(c) KCI and NaCt solutions possess sems vapour pressure.
(d) KCI solution possess sems vapour pressure.
(d) KCI solution possesses lower treasing point than NaCl solution.

The molal bettleg point constant is the ratio of the elevation in botting point to the fraction of solution (d) mola fraction of solution (e) molarity (b) molarity (c) mola fraction of solution (d) molarity (d) molarity (e) mo



Colligative properties are the properties of

- (a) Dilute solutions which behave as nearly ideal solutions
- Concentrated solutions which behave as nearly non-ideal solutions
- (c) Both (a) and (b)

.d) Neither (a) nor (b)
opur Board, 2009) (Roundplood Board, 2009)

ASSET RS TO MULTIPLE CHARL QUESTIONS

it Ans et

Consider draft of water 1 draft drafter = 1 kg = 1000 g Molecular mater of water = 18 g thor is

Number of moles of water to 1 dro² = $\frac{1000}{10}$

=55.55 moto

State 55.55 (woter of witter are present in 1 dm², therefore, moladity of pure winer is \$5.65

Mass of glucose = 75 g Molecular mass of plucose = 180 d

No. of moles or glucose = $\frac{18}{180}$ = 0.1 moles

Mass of water = 90 g Meteoday mass of water = 18 g

No. of moles of water= 16 = 5 moles

Thus mole fraction of glucooz= x_0 = $\begin{bmatrix} 0.1 \\ 5 & 0.1 \end{bmatrix}$ = $\begin{bmatrix} 5 \\ 1 \end{bmatrix}$

According to Regult's law

E.g. maler of glucous ≈ 180 g. 10% solution means 10 g. of glucose and present to 100 cm² of solution. Thus: 100 cm² of solution. Thus: 100 cm² of solution. Thus: 100 g. of glucose will be present in solution ≈ 100 gm² 180 g. of glucose will be present in solution: 100 ±150 g. of glucose will be present in solution: 100 ±150 g. of glucose will be present in solution:

er Aus. (b)

It as a successful mixture of two liquids bods at source temporabute than either of them, then it mixture to account or pressure of the mixture is more than the ideal vapour pressure. Hence, this mixture well show positive deviations from Roouli's law.

wiit Ans: (a)

CET Ans: (a)

Envision in boiling point is a collegative paperty and is desertly proportional to the number of purifice in the solution.

5.85% NeCL 18% glacose and 6.0% uses rotutions are all I hotels solutions. However, NeCL innices in cooler to give double number of from Date to greater humber of perificial in NeCL solution, its boiling point will be hartest.

Assumer of person
will be haptest
tixt Ann (b)
Elevation in boiling point a given or

$$\Delta T_{k} = |a_{k}| \times m$$
 or $\frac{\Delta T_{k}}{m} = |k_{k}|$

Hence, model beliefly point constant (bb) is the ratio of the absolute in boding point (Δ Tb) to the modelity of solubon (m).

 $\frac{\Delta P}{p^2} = a_k \approx \frac{1}{51}$

Ethanol and water for "in a non-ideal solution. This solution shows positive as: ladons from Reputify law. Hence vepolur pressure of this solution will be more than their of water.

10-70-10

In the formation of solution that shows positive deviation from Recult's law (coops of altractions among motivation are decreased, Therefore motivation and properties and present away from each other Hames, volume of phillion is increased.

Both NeCl and KCI contest in water completely in give double marber of lines when some moles of NeCl and KCI for when some moles of NeCl and KCI for dissolved separately in fixed amount of water. However, they produce sums number of lent in water. However, they be the some some however are values of colleging properties like veryout pressure towering at:

(xl Aust fa)

in duty solutions, particles do not have torces of administration artenders around themselves. Therefore, the solutions behave ideally and collegative properties are observed.

faller Chemietra, Pere-I

541 gif in the blanks with suitable words

Number of molecules in 1 dm³ of 1 M sugar solution is Ŋ 100 g of a 10% ag. solution of NaCH contains 10 g of NaCH in

When an accordance is distilled, its temains constant.

The motal freezing point constant is also known as constant.

The boiling point of an azeotropic solution of two equids is lower than either of them because the solution shows

them because the solution shows from Raque's law.

Among equimoral ag solutions of NeCl, BaCl₂ and FeCl₃, the maximum depression in freezing point is shown by _____ solution
A solution of athanol in water shows

deviations and gives assotropic solution with ____ boiling point then other components,

Colligative properties are used to calculate of a compound

The hydration energy of Br ion is then F on The agreeus solution of NH₄Cl is while that of Na₅SO₄ is

(i) position deviations (ii) less (x) acidic neutral (iii) less (x) acidic neutral



Indicate True or False from the given statements

At a definite temperature, the amount of a solute in a saturated solution is found

Polar solvents readily dissolve non-polar covalent compounds.

The molecular mass of an electrolyte determined by lowering of vapour pressure as less then the theoretical molecular mass.

The rate of evaporation of a Equid is inversely proportional to the intermolecular forces of attraction

The solubility of a substance decreases with increase in Temperature if the head of

a solution is negative. The rate of evaporation of a liquid is inversely proportional to the intermolecular

forces of attraction (M) All solutions containing 1 g of non-volatile, non-electrolyte solutes in same solvent

will have the same frazzing point.

Will The frazzing point of a 0.05 molal solution of a non-electrolyte in water is -0.93

Hydration and hydrolysis are different process for Na₂SO₄

The hydration energy of an ion only depends upon its charge

Trop (ii) False (iii) Trop (ic) Trus (clii) Fulse (lis) False (viii) False



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Charles Park Define and explain the followings with one example in each con-

(b) Zaotropic solutions

(d) final of hydration

(c) A estimation of solid in a polid

(i) Water of crystallization

igi A consulate temperature

(h) Assotropic solution

(t) A non-trient solution

(j) Conjugate solution

Solved in the chapter

Q6 (a) What are the concentration units of solutions. Compare moler and ad-

Solved on Page 504

(b) One has one molel solution of NaCl and one molel solution of glucose.

(f) Which solution has greater number of particles of solute?

NaCl dissociates in H₂O, while glucose is not ionized. NaCl produces Na* and Cl*iss. Thus number of perticles are increased. Hence, 1 motal NaCl solution contain gratering. number of particles than I motal glucose solution.

(ii) Which solution has greater amount of the solvent

Both are 1 motel solution i.e. 1 mole of solute is dissolved in 1 kg of solvent. Therein both will have same amount of solvent.

(iii) How do use concert these concentrations into weight by weight percentage?

For Emistal Safet schutzen	Tor Landal alla age solution
Moles of NeCl = 1	Mass of glucose dissolved = 180 g
Here of NoCi - \$8.5 g	Mass oil solvent (water) = 1000 g
Stree 1 mole of NaCl is dissolved in 1 by of	Potes man of solution = 1000 + 180
solverst, therefore	= 1180 g
Man of solvent (water) = 1000 g	Total mass of glucose x100
Total remain of the solution = 1000 + 58.5	Total mass of solution
= 1056.5	% of glucose = 180 x 100 x 15.2%
% of ReC= Mass of NeCl Total mass of solution > IIII	1180
% of PMC1= 98.5 .056.5 = 100 = 1.059	

October Paris

Explain the following with r

The concetration in terms of molester is indepen depends upon temperature.

Japanen Board, 2007 Gujranuculo Board, 2009: Falsolchod Board, 2007, 2011 Multon Board, 2011; at life the concentration of a molar solution is changed by unriging of temperature

@.G. Johan Round, 2000)

Molality is defined as the number of moles of solute dissolved in 1 kg of solvent, while Molarity is defined as the number of males of solute dissolved in 1 dm³ of solution.

Since molestity is based upon mass, therefore, it is independent of temperature. While polarity is based upon volume, therefore, it varies with temperature due to change in

II) The sum of male fractions of all the components is charge equal to unity for any

|Falsafabad Board, 2008, 2013: Latters Board, 2010: D.G. What Sound, 2012: Hulton Board, 2012) If Press that sum of male fractions of all components is always equal to tine, (Paudpini Board, 2008)

Consider a solution of two components A and B

Let motes of A are na and that of B are na

Their mote fractions are given by

$$x_A = \frac{n_A}{n_A + n_B}$$
 _____, 1 | $x_B = \frac{n_B}{n_A + n_B}$ _____ 2 |

Their sum of mole fractions will be

$$x_A + x_B = (3)$$

Put eq (1) and (2, in eq. 3)

$$\frac{n_A}{n_A + n_B} + \frac{n_B}{n_A + n_B}$$

$$\frac{n_A + n_B}{n_A + n_B} = 1$$

 $n_A + n_B$ Hence, sum of mole fractions of different components in a solution is equal to one.

100 # of 98% H SO, how a polume of 54.88 cm 98% H_aSO_a has a density = d = 1.94 g/cm³

Mass of $M_2SO_4 = m = 100 g$

Volume of H₂SO₆ = V = ?



niero Parisi

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 $d=\frac{m}{V}$

or
$$V = \frac{m}{d} = \frac{100}{1.84} = 54.38 \text{ cm}^3$$

lelective lowering of vapour pressure is independent of the temperature.
(Panalobul Board, 2006 Sergodio Board, 2009 Hazolpind Board, 2009: Guironacio Roard, 2019. According to Reput's law

Relative lowering of vapour pressure is equal to the mole fraction of solute

Where $\frac{\Delta P}{n^0}$ = Relative lowering of vapour pressure

While
$$x_2 = \frac{h_2}{n_1 + n_2} = \text{mole fraction of solute}$$

Since number of moles are not changed with temperature. Hence, relative lowers vapour pressure is independent of temperature

Calligative properties are obeyed when the solute is non-electrolyte.

Collegeave properties depend upon number of solute particles. If solute is electrical en it will nonize and change the number of particles in the solution. Hence, elected colligative properties will be observed

(eii) Colligation properties are obeyed when the solutions are dilute Off Colligative properties are obeyed when the solute is non-electrolyte and also when the solution are dilute Explain with reason. (Currenteels Board, 2009: Rountpind Board, 2012; Makes Board, 2014).

Collegative properties also depend on the fact that solution should be dilute to that en te particle behave ardependently

in conc. solution, solute particles don't behave independently. They have attack each other. Thus, normal colligative properties are only observed when the solution is did

(all) The total volume of the solution by mixing 100 cm² of conter with 100 cm² alcohol may not be egood to 200 cm² disselfs it.

Duning ideal volumes formation and solution is a set of the solution formation and solution is a set of the solution of the solution of the solution is a set of the solution of the solution of the solution is a set of the solution of th

During ideal solution formation volume is not changed. However, Alcohol and all is not changed. However, Alcohol and all is not changed. form a non-detail solution. Forces of attraction between water and alcohol in pure state of greater than in solution. There is a struction between water and alcohol in pure state of greater than in solution. grapher than in solution. Thus, it shows positive deviation from Racults law. During a formation of this solution heat is when their control of the solution heat is when their

Hence 100 cm² of H₂O and 100 cm² of alcohol may not be equal to 200 cm²

(mill) One molel solution of urso in water is dilute as compared to one moler volution area, but the number of particles of the solute is some. Justify it (Faladonal Boord, 2017, 2010; Acad Kanbrit Boord, 2017, 2

other Characters, Part-1

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of One model glucouse solution to diffuse in compared to one moles solution beard. 2008: Faisatabad Beard. 2009: Resulpted Board. 2011. Copins for 1, 2013. Surgodin Board. 2014.

Both 1 motal and 1 motar urea solution contains i mote of urea. Hence both these philipps contain same number of particles i.e. 6.02 x 10m

For 1 molel solution, 1 mole of trea is dissolved in 1 kg of water up 1 day However,

For 1 molar solution, 1 mole of usea is first taken to a measuring flask and then volume smade up to 1 dm³

Hence, 1 molar solution contain less amount of H₂O than 3 molal solution.

qr. What are non-ideal solutions? Discuss that types and give three example of each.

(a) Explain fractional distillation dustify the had causes when coplotted against boiling point of solutions.

b) The solutions should positive and regains desinious cannot be fraction et their specific compositions. Explain it.

Q9. (d) What are executrapic mistures? Explain them with the help of graphs? Of West are executrapes for executrapic misture? Give one example. D.C. Kom Bourd, 2007: Multan Board, 2009: Labore Board, 2016. Education Board, 2017: Sugardia B. 2019.

312, 2014)

M Explain the effect of temperature on phenol-union epitem.
School on Page 514

910. (a) What are the colligative properties? Why are the celled so?
(B.G. Khan Board, 2007: Lohars Board, 2009: Makes Board, 2010: Roses

Oil Why some properties are called collegated?

(Surporth Board, 2007 D.S. (Dan Board, 2011 Behavelor Board, 2011)

(Surporth Board, 2000 D.G. (Dan Board, 2011) Behavelor Board, 2011)

lobed on Page 524

What to the planeted atgrafficance of R₂ and R₃ selam of actions of solutions of actions of solutions of

Hose do you explain that the locaring of expour presents is a collective property!

Hose do you explain that the locaring of a non-volatile, non-electrolyte solute in a polarie. polatile eolpent?

bord on Page 625

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Que How do you justify that

(a) Boiling points of the solvents increase due to the presence of solutes
(Fassiahad Board, 2017, Gujranunia Board, 2010, Multan Board, 2013, Lujan

Why the boiling points of a solution of a non-volatile solute in a volatile always greater than the boiling point of pure solvens?

Burd 2008,

When a non-vocable, non-electrolyte solute is dissolved in volatile solvent, its on pressure is lowered. Thus to boil solution more heat is required to bring its up

pressure equa, to external pressure. Hence, boiling point of solution is increased. (b) Freezing points are depressed due to the presence of solutes (Lahore Boord, 2007-2009). Faisaluban Board, 2010. 2011. Rawalpindi Board, 2011. Guraninto

OR Why the freezing point of the solution is always less than the freezing point of

When a non-volatile, non-electrolyte solute is dissolved in volatile solvent, its ran pressure is lowered. Due to lowering of vapour pressure, solution freezes belo freezing point of pure solvent. Hence, freezing point of solution is depressed.

(c) The boiling point of one molal were solution is 100.52 °C but the boiling p of two molal were solution is less than 101.04 °C.

Elevation of boiling point is a colligative property. Norma, colligative properties observed when the solution is dilute so that each solute particle behave independently

Two modal area solution is cone than one mail solution. In concentrated two particles particles do not behave independently They have attractions to other Hence one modal solution boils at 100.52°C. While two molal solution boils at 100.52°C. abnormally below expected at 101 04°C

(d) Beckmann thermometer is used to note the depression in freezing point.

(Rawaipind: Board, 2007: Multan Board, 2009: Labore Board, 2013. Bahasaipin B 201.)

There is very small difference between freezing point of pure solvent and is discussion. Ordinary thermometer can read upto 0.5 K (upto one deems) place) these cannot differentiate between freezing point of pure solvent and solution. Beckmann thermometer can read upto 0.01 K (upto two decimal places), there can exactly measure the freezing point of pure solvent and solution.

(e) In summer the antifreeze solutions protect the liquid of the radiator boiling over boiling over. Bourd, 2007)

Antifreeze solution consists of solution of ethylene glycol in HaO Since et and is non-volatile therefore giveol is non-volatile, therefore, vapour pressure of this solution is lowered and to point is increased. point is increased.

In summer, due to large heat in engine, HaO may boil over in radiator. How

Chemistry: Part-1

547 to were vapour pressure of solution, it requires much greater heat to boil. Hence, this

moders prevents room and to lower the melting point of ice. (Surjedia Board, room Gajranwala Board, 2016, Antien Board, 2013, Surjedia Board, 2013, Multan Board, 2013, Bullian Back or KNO₃ is added to H₂O, its vapour pressure is lowered. Due to lowering d support pressure, solution freezes below the freezing point of H_eO. so it will have more cooling effect.

Hence, this mixture is used as freezing mixture e.g. in an ice-cream machine

What is Rapull's law. Give its three statements, How this law can help us to understand the ideality of a solution,

Solved on Page 515

Give graphical explanation for elevation of boiling point of a solution. Describe one method to determine the boiling point elevation of a solution.

Solved on Page 527

Preezing points of solutions are depressed when non-volatile solutes are present in volatile solvents. Justify it. Plot a graph to elaborate year answer. Also, give one method to record the depression of freezing point of Dr#. a solution

Solved by Page 530

Discuss the energetics of solution Justify the heats of solutions as exothermic and endothermic properties.

Solved on Page 533



College Chapters: Part-I

% by weight

Molarity

$$M = \frac{No \text{ of moles of solute}}{Volume of solution in dm3}$$

$$m = \frac{No \text{ of moles of solute}}{\text{Weight of solvent in kg}}$$

$$m = \frac{wt \text{ of solute}}{\text{Maleculer wt of solute}} \times \frac{1}{\text{Weight of solvent in kg}}$$

$$\mathbf{x} = \frac{\text{No. of moles of one component}}{\text{Total no. of moles of all components of solution}}$$

Perts Par Million (ppm)

$$P = P^{e} \, \kappa_{1} \quad \text{or} \quad \Delta P = P^{e} \, \kappa_{0} \, \phi_{1} \cdot \frac{\Delta P}{P^{0}} = \kappa_{2}$$

Lowering of V.P./Moi. Mass determination etc.

$$\frac{\Delta P}{P^0} = \frac{w_2 M_s}{w_1 M_2} \quad \text{or} \quad M_2 = \frac{P^0}{\Delta P} \times \frac{w_2 M_1}{w_3}$$

Elevation of B.P./ Mol. Mass determination etc.

$$\begin{array}{lll} \Delta T_b = k_b \times m & \text{for} & \Delta T_b = k_b \times \frac{max}{M_Z} \times \frac{1000}{w_0} & \text{or} & M_Z = \frac{k_b}{\Delta T_b} \times \frac{w_2 \times 1000}{w_1} \\ \text{and Boiling point of solution} = \text{Solling point of solvent} + \Delta T_b \end{array}$$

Depression of F.P./ Mol. Mass determination etc.

$$\Delta T_1 = y_0 = m_0 \quad \text{or} \quad \Delta T_2 = y_0 \times \frac{M\Delta}{M_2} \times \frac{1000}{w_1} \quad \text{or} \quad \frac{y_0}{\Delta T_1} \times \frac{y_0 \times 1000}{w_1}$$
 and Financing point of solution = Financing point of solvent — ΔT_1

NOMERICAL PROBLEMS TREM 1SI

(a) Calculate the molerity of plucose solution who q.17 250 cm² of solution.

Court on:

Mass of glucose
$$= y = 9g$$

Mal. Mass of glucose $= M = 180g$

Volume of solution
$$\approx V = 250 \text{ cm}^3 \approx \frac{250}{1000} \text{ cm}^3 = 0.25 \text{ dm}^3$$

Molarity of solution = 7

Molarity of solution is given by

Molarity =
$$\frac{w}{M} \times \frac{1}{V(dm^5)}$$

Motarity =
$$\frac{9}{180} \times \frac{1}{0.25} = \boxed{0.2 \text{ mol / dm}^3}$$

(b) Calculate the mass of ures in 100 g of water in 0.5 motel

Mass of water solvent $\}$ = W = 100 g = $\frac{100}{1000}$ = 0.1 kg

Molality of solution = m = 0.3 Motar mass of urea = M = 60 g/mol Mass of area dissolved = w = 7

Molality is given by

$$m = \frac{w}{M} \times \frac{1}{W(kg)}$$

$$0.3 = \frac{w}{60} \times \frac{1}{0.1}$$

$$w = 0.3 \times 60 \times 0.1 = 2.8 \text{ g}$$

(d) Calculate the cano, of a schutton in motally which is obtained by mixing 250 g of 20% solution of NaCl. deline

250 g of solution contain NeCt =
$$\frac{20}{100} \times 250 = 50$$
 g.

100 g of solution contain NaCi = 40 g 200 g of solution contain NeCi = $\frac{40}{100}$ × 200 = 80 g. Hapen after mixine Total mass of NaCl = w = 50 + 80 = 130 gFormula mass of NaCl = M = 58.5 g/mol Mass of total solution = 200 + 250 = 450 g. Mass of Water = W = 450-130 = 320 g = 0.32 kg. Moiality = m = ? Molality is given by $m = \frac{w}{M} \times \frac{1}{W(kg)}$ $m = \frac{130}{58.5} \times \frac{1}{0.32} = 6.94 \text{ m}$ ġ Q.18 (a) An eq. solution of sucrose has been labelled as one motel. Find the not freation of solute and the salment. Bahitlan 1 motel no sucrose solution means that 1 mole of sucrose (342 g) has been dissolved in 1 kg ($1000~{\rm g}$) of H_gO . Mance Moles of storoes = n_{me} = 1 mole Mass of H₂O = 1000 g Mol Mass of H₂O = 18 g/mol Moles of $H_8\Omega \approx n_{\text{NSO}} \approx \frac{1000}{18} = 55.55 \text{ moles}$ $x_{\text{time}} \approx \frac{1}{1 + 85.55} = 0.9177$

(b) You are provided with \$0% sulphurto and having density 1.8 giam! How much volume of this sulphurta acid sample is required to obtain 1 dm² of 20% HyBo-tehace density is 1.25 giam!

= 1.8 g

 $\chi_{\text{eqs}} = \frac{55.85}{1 + 55.55} = 0.9883$

% of sulphuric acid = 80 % Density of sulphuric acid = 1.8 g/cm²

940

College Characters: Part d

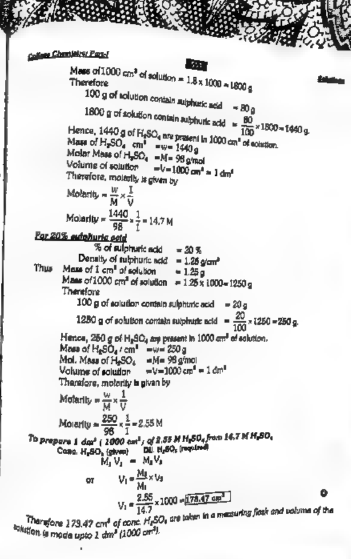
For F" solution

Solution

Cor 80%

ticinitaria ecid

Thus Mass of 1 cm³ of solution



al ent William

ø

Solution:

1 mole of K₂SO₄ produce 2 moles of K* ions

Therefore, 0.2 moles of K₂SO₄ produce 0.4 moles of K* ions

Hence conc. of K* ions = 0.4 moles/dm⁴

1 male of KCI produces 1 moles of K* ions. Therefore, 0.2 moles of KCI produces 0.2 moles of KC ions. Hence, conc. of K* ions = 0.2 moles/dm².

After relating two moles solutions. Conc. of K^* tons = 0.2 + 0.4 = 0.6 moles in 2 dm^* if 250 cm^* of each solution is mixed then

Total Volume = $250 + 250 = 500 \text{ cm}^6 = \frac{500}{1000} = 0.5 \text{ dm}^6$ Hence 2 dm⁶ of solution contain K* tons = 0.6 moles

0.5 dm³ of solution contain K* ions = $\frac{0.6 \times 0.6}{2}$ = 0.15 moles/dm³

1 dm² of solution contain K' ions = 0.3 moles/dm²

Q.20 5 g of NaCl is dissolved in 1999 g of water. The density of resulting solution is 0.997 glant. Calculate the molality, molarity and male fractions of this solution.

Assume that the solution is equal to that of the solvent.

Kololite

Mass of NaCl = w = 5 g Formula mass of NaCl = M = 58.8 g/mol Mass of H₂O = W = 1000 g ≈ 1 kg

 $\begin{aligned} & \text{mobility} = \frac{w}{M} \times \frac{1}{W} \\ & \text{mobility} = \frac{5}{58.8} \times \frac{1}{1} = \boxed{0.0854 \text{ m}} \end{aligned}$

Melecity

Mass of NeCl = w = 5 g Formula mass of NeCl = M = 58.8 g/mol Mass of H_gO ≈ W = 1000 g = 1 kg

Since volume of solution is equal to the volume of solvent. And 1 kg of water $= 1 \text{ dm}^3$. Therefore volume of solution $= V = 1 \text{ dm}^3$.

Giben Chambelon: Part-j

Molarity is given by

molarity =
$$\frac{w}{M} \times \frac{1}{V}$$

molarity = $\frac{5}{58.8} \times \frac{1}{1} = \boxed{0.0854 M}$

Mole

Mass of NaCl =
$$w = 5$$
 g
Portoule mass of NaCl = $M = 58.8$ g/mol
Moles of NaCl = $\frac{5}{58.5} = 0.0854$ moles

$$\begin{array}{ll} \mbox{Mass of H_2O} & = W = 1000 \ \mbox{g} \\ \mbox{Mol. Mass of H_2O} & = M = 18 \ \mbox{g/mol} \\ \mbox{Moles of H_0O} & = \frac{1000}{18} = 55.56 \ \mbox{moles} \end{array}$$

Hence

$$\times_{\text{N=O}} = \frac{0.0854}{0.0854 + 55.56} = \boxed{0.00154}$$

$$\chi_{\rm H2O} = \frac{55.56}{0.0854 + 55.56} = \boxed{0.9984}$$

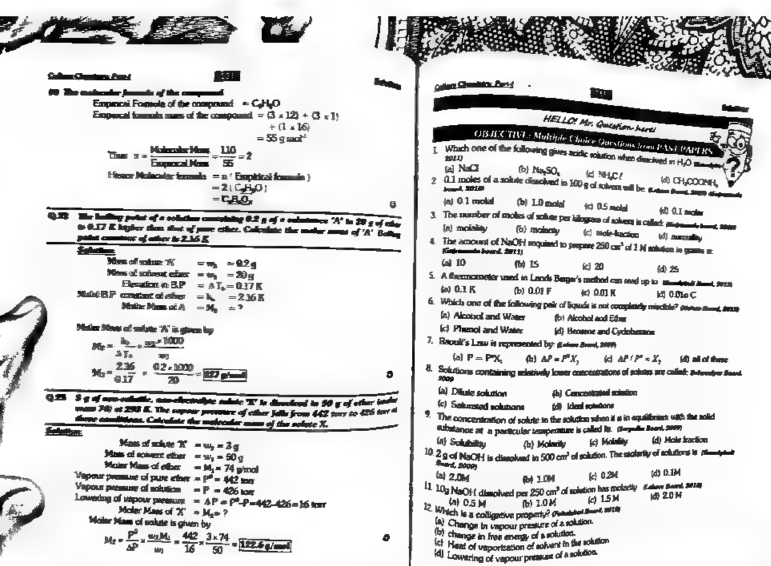
Q.21 4.675 g of α compound with captrical formula C_pH₁O, was discolored in 212.5 g of pure homeone. The finating point of the solution was found 1.02°C less than that of homeone. The motor fracting point constant of benzens to 5.1°C. Calculate.

() The relative motor must of the compound

Mass of solute compound $= w_z = 4.675 \, \mathrm{g}$ Mass of solute compound $= w_z = 4.675 \, \mathrm{g}$ Mass of solutent benzens $= w_1 = 212.5 \, \mathrm{g}$ Freezing point depression $= \Delta T = 1.02$ Motal F.P. constant of benzene $= k_1 = 3.1^{\circ}\mathrm{C}$ Molecular Mass of compound $= M_2 = 7$ Molecular Mass of the compound is given by $M_0 = \frac{k_1}{M_1} \times \frac{k_2}{M_2} \times 1000$

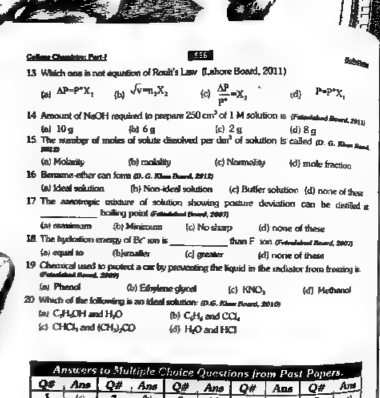
taker Masse of the compound is given by
$$M_{R} = \frac{k_{T}}{\Delta T_{r}} \times \frac{v_{Z} \times 1000}{\omega t}$$

$$M_{R} = \frac{5.1}{1.02} \times \frac{4.675 \times 1000}{212.5} = \boxed{110 \text{ g/rssel}}$$



V Jahar

THE STREET



Γ	D	etatle	d Ev	alonas	don .	(D.	-	7.0	000	2.
		_								
-	1100	(a)	17	(b)_	18	(c)	19	(b)	20	<u> (b)</u>
ł	11	(b)	12	(d)	13	(b)	14	(a)	15	
4	4.6	(6E)	7_	(4)	8	(a)	9	(a)	10	(4)
- 1	-	1-7		- 907		129	4	(20)	2_	

xplanation of Past Papers MCQs & answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK! College Chemisters: Fort-1

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SHORT & LONG OVESTIONS FROM PAST PAPERS CONCENTRATION UNITS OF SOLUTIONS

- that do you mean by percentage talls composition of a solution, our (2) Define motority. Give one example (Matter Source, 2014)
- (3) What is molality. Give its formula? (Fundame Board, 2016)
- (3) What is recovery. Since as partitles. We had a sound 1909 Labore Board, 19031
 (4) Define molarity and malify. Gaue their mathematical expression. (Labore Energy, 1904)
 (5) Define mole fraction and parts per million, (Sargodia Board, 2019)
- (6) What is (or Define) parts per million (ppm). Give its mathematical expression middle Board, 2007 Labore Sound, 2009; D.G. Rheer Scend, 2009; Oal Define parts per million and when this concentration unit is used. (Supplie Seed, 201) on ungue parts this concentration unit is used. (Supplie Seed, 201) (1) How will you prepare 0.2 M NaOH solution? (0.5, 15th Beerl, 1000)
- (5) Flow will you prepare 5% W/V used solution in takes the state south seems. In
- (9) How will you prepare 10% W/V wea solution in uniter? Triangled Board, 2019.
- (10) Calculate the percentage by weight of NaCl #2g of k is dissolved in 20g of unter (Bespecific Board, 2011 Lateurs Board, 2010)
- (Buryothe Board, 2011 Laters Board, 2010)

 (11) What is the molelity of a solution prepared by dissolving by of glucose in 250g of Citales (Surpodia Board, 2013)
- (12) Find the molarity of a solution containing 41.4 g fo K₂CO₃ dissolved in 500cm³ of given solution. Molar mass of K₂CO₂ is 138 g/mol. others is
- Calculate the molarity of a solution containing 20.7g of K₂CO₃ dissolved in 500 cm³ of the given solution. (D.G. River Secret, 2014)

- lana Overstona

 (ii) Define the following terms with examples.

 (ii) Molarity (iii) Mosality (iii) Mole fraction (iv) Parts per million awardles bood, at laters Beant, 2016;

LIQUID-LIQUID SOLUTIONS, NADULTS LAW

ers Overstone

- (1) What are confugate solutions? Give examples, (taken bear), 2009.
- What is consulate temperature or critical solution temperature? Give example phases found, 2007 to G. Niew Board, 2017 to G. Niew Board,
- What is Racults into ? (Rendyles Bord, 1915)
 (4) Cane sugar cornol be dissolved in benzene. Give reason, (Repeated Bord, 1915)
- (5) Why glucose is soluble in water but insoluble in CCL? (harmone Board, 1869)

(ii) State one form of Floorit's law. (iii) pages floors, 2010) (2) (as a part of question) (2) State one form of Rocult's law, (Reduced Beard, 2010) (2) has a part of global a law? Give State different forms of Rocult's Law, (Letters Beard, 2011) on What is Rocult's Law? Give the identity of a solution. to time statements. How this kno can help us to understand the ideality of a solution. (2.4. Rise Board, 2015) On Give three statements of Results loss and the mention hou



College Chapters: Part C

655

Repulse low helps in determining the ideality of a solution, they Secret. 2017 44

(3) Define Raoult's into and derive the equation AP = P * X_q (Surpolite Its) end. 2014b

(4) What is Rapult's law? Explain when both components are volatile (Muller Bo

(5) Discuss Roults Law for the solution in which both component are volatile. (Ampedia). MALE Federal and Second, 2013)

EDEAL AND HOW-IDEAL SOLUTIONS

- (1) Give any two points which show the ideality of a solution. (Labora Board, 2008)
- (2) Give two pairs of liquids which form ideal solution. (D.G. Khan Board, 2015)
- [3] Differentiate between ideal and non-ideal solutions, (Bahmalaur Board, 2009 sheed, 2011 Labors Board, 2015)
- (4) Non-ideal solutions do not obey Roculé's law Explain (Surgethe Sweet, 2009; Result Speed, 2010)
- (5) Many solutions do not behave ideally. Give reason, (Gyrennia Bond, 2014)
- (1) Explain the difference between ideal and non ideal solution. (6-1) miles Bound, 2012) OR Distinguish between Ideal and non-ideal solution. (Behavior See (2010) (2) (as a part)

ZEOTROPIC AND AZEOTROPIC MUXTURES, DEVIATIONS FROM RADULT'S LAW Short Overflone

- (1) Define Zeotropic mixtures. Give one example. (Laiore Bond, 2014)
- (2) What is positive deviation from Raoult's taw? (Letter Round, 2007)
- (3) Differentiate between zeotropic and assotrapic mixtures. (Mutter Board, 2011. Lena Questions
- (1) What are non-ideal solutions (or agealrapse)? Discuss their types with example (Fateshed Board, 2007: Original Board, 2013)

 (2) Define non-ideal solutions and explain positive decision with the help of a graph.

 (Benedictal Board, 2014)
- (Bergreike Board, 2013)
- (3) Explain negative deviations from Repult's law. (0:6. Khee Board, \$6) ()
- (4) What are idea solutions? Graphically explain that how the ideal solutions of two liquid can be separated by fractional distillation. (D.G. Khen Board, 2009)
- (5) What are toled solutions? Expirin the fractional distillation of ideal mixture of two liquids. (Labora Board, 34(2))
- (6) Explain the fractional distillation, dustify the two curves when composition is plotted assistant believe materials of the curves when composition is probable. against boiling point of the solution. (Februaries Search, 2012)

SOLUBILITY SOLUBILITY CURVES, PRACTICINAL CHYSTALLIZATION Short Questions

- (1) Define solubility, curve. Name its two types, (Felorished Board, 2019)
- (2) Why some substances show discontinuous curves? (0.6. Khen Board, 2004)
- (3) Differentiate between continuous and discontinuous solubility curves. (Letters Seet 481)

Charleton Carri

17.5 (i) Why fractional crystalization is carried out? (the this fractional crystalization House is under the Band 1916 But of Define fractional crystalization House is under the Band 1916 But of 1921

Described in Opine solubility curves and gives in types with suitable examples, or a countries and make class different stable examples. Define solubility and make clear difference between continuous and discontinuous solubility curves. (Faterland Board, 3916: Caparanala Board, 3911).

DUBATIVE PROPERTIES

get Questions

| Define colligative properties. Nome some important colligative properties. 49.0. Sec. 5010: Labour Board, 2011; Februard Board, 2018; Samuelle Board, 2016.

p Deline ebuiltoscopic constant for molal elevation constant or molal bailing point constant) with one example (Leben Book Loss)

I) What is molel freezing point constant for cyroscopic constant) (Indicated in What is cryoscopic constant. Give its value of water (tempeths Beerl, 2007)

il) State Ebullioscopic constant and Cryoscopic constant. (Matter Based, 40) D 5) Give the conditions of colligative properties. (Europh South 2011)

ii) What are colligative properties? Closely these and explain any one throughout the

ORERING OF VAPOUR PRESSUES, SLEVATION OF BOILING POINT DEPRESSION OF REZINO POINT, APPLICATIONS OF COLLIGATIVE PROPERTIES

Ret Cassions

While is the vopour pressure of a solution less than pure solvent? (Line Sound, 2014)

While is the vopour pressure of a solution less than pure solvent? (Line Sound, 2014)

Depression in freezing point is a colligative property Justify (Labor Board, 2013)

Gue too applications of collegative properties (Samuella Board, 8913).

Etylenes glycol may be used in radiator of an outsmobile. Give too aspects of its use feeling Board, 2008. Guyennade Board, 2019.

Define collegations

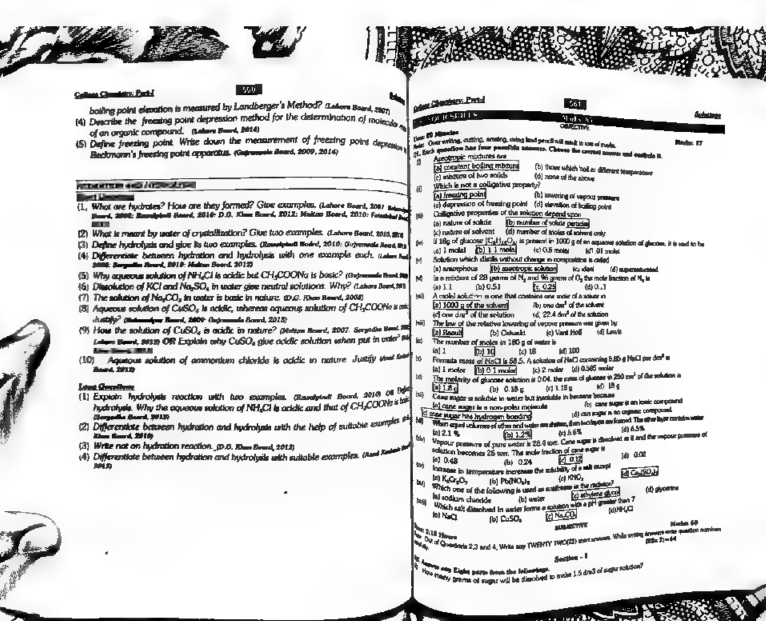
Define collegative properties. Describe the lowering of uppour pressure in terms of collegations.

colligative property of solution and derive the expression for molecular mose of non-

tolgille solute. (Schowelper Seers, 1929) Gibbs graphical explanation for elevation of bailing point of a solution. (Leheve in the control explanation for elevation of bailing point of a solution.) Prophical explanation for eteration of boiling point of a science have, area on Define of Expiral advantion of boiling point with the help of figure. (Anhore have, area on Expiral believation of boiling point. (Familiable Beard, 2011) on Expiral between of boiling point of a solvent by solve with the help of a graph. Derive formula for determination of molecular mass of an unknown solve from this concept. (D.B. Beard, 2011) on the help of figure. (Making Beard, 2011) on the help of figure.

*** A series of molecular mass of an unknown source from this concept, the series are series of the unit of Balling Point with the help of figure. White the source of Balling Point and Basel, 2011 \$412. Behavedow Board, 2011 disjourned Basel, 2010 to determine the boiling point of unfolding point elevation. Describe a method to determine the boiling point and four if an objection. (Sempotine Board, 800) of What is the elevation of boiling point and help if can be measured by Landberger's method? **Resemble board, 2007 Gujernseals and 2012; **Sempotine Board of Define and classify colligative properties. Flow * can be measured by Landberger's memoor measure properties. How 2012; Amenda Band 2011 on Define and classify collegative properties. How

and the same of



EVERYORAL 1

COLLEGE COLLEGE CHEMISTRY

Intermediate Part I





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Tallat Jamshed

Zaheer Abdullah

Shahid Ali

M. Ashraf Kamal

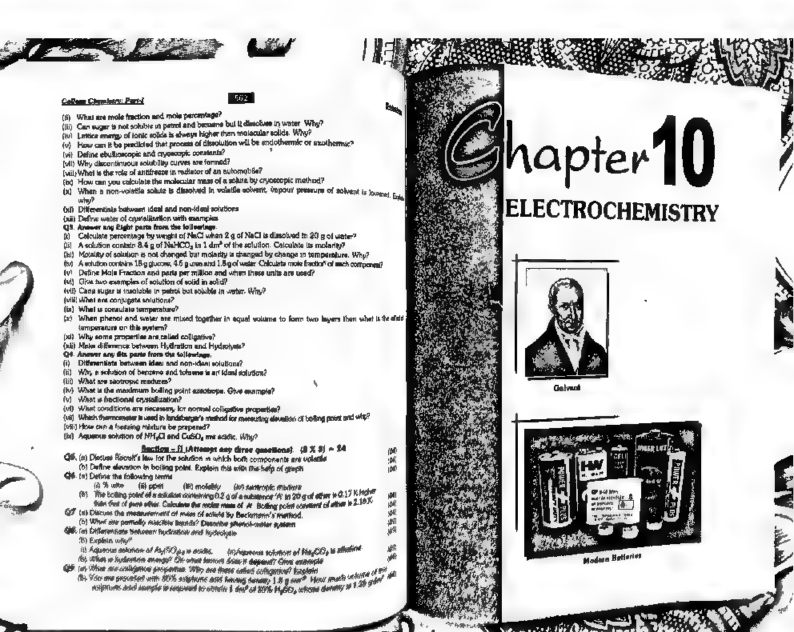
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🦟 🐣 CONTENTS 🐗

Chapter-10 Walter to the Manager

Electrochemistry

OXIDATION STATE AND BALANCING OF REDOX EQUATIONS

Oxidation number or exidation state Balancing of equations by oxidation number method

Balancing of equations by ion-electron method

ELECTROLYTIC CONDUCTION

Electrochemical cella Electrolytic cells

Explanation of electrolyals

Electrolysis processes of industrial importance

Voltare or Galvanic cell(Zn-Cu cell)

ELECTRODE POENTIAL

Standard hydrogen electrode (SHE)

Measurement of electrode potential

THE ELECTROCHEMICAL SERIES

Applications of electrochemical series

MODERN BATTERIES AND FUEL CELLS

Lead accumulator (rechargeable)

Alka ine battery (non-rechargeable)

Silver oxide battery

Nickel cadmium cell (rechargeable)

Fuel cells

Objective and short answer, questions (exercise)

Numerical problems (exercise)
Past Papers MCQs and Short Questions

Test your skills

ter Chamberr Part-I

MRODUCTION

Electrochemistry

The branch of chemistry, which deals with the conversion of electrical energy into shall energy and chemical energy into electrical energy, is called electrochemistry.

Sign

Orldetton

it may be defined as the addition of anygen, loss of hydrogen or loss of electrons from

Peduction

it may be defined as the removal of anygen, addition of hydrogen or addition of ectrons from a species.

Addition Reaction

A reaction to which a substance losse electrons is called axidation reaction.

e.g.
$$Z_{D_{(0)}} \rightarrow Z_{D^{(4)}} + 2e$$

duction Reaction

The reaction in which a substance gains electrons is called reduction reaction.

Oxidation reduction reactions are also known as redax reactions.

in a reduction, exidation number of alements undergoes changes

AND THOSE AND BALANCING OF REDOX EQUATIONS

WALLER OF COLUMN DAYS AND A STATE OF apparent charge on an atom of an element in a molecule or an ion.

they be positive or negative or refe.



College Chambian, Part-I



- The oxidation state of free H₂ is zero
- The oxidation state of Na and Cl are +1 and -1 respectively

Rales for Assigning Osldation Nanders

- The oxidation number of a free element is zero, e.g. H₂°, Na°, Mg°
- 2. The oxidation number of hydrogen is +1, except in the case of metal hydrogen NaH, MgH, where it is --1
- 3. The excidation number of except is 2, except in the case of metal peroxides, when it is -1, in superoxides, where it is -1/2 and in OF, where it is +2.
- 4. In binary compounds, the exidation numbers of group VII A halogens is 1.
- 5. The exidation number of each element of the groups I-A, I.-A and III-A are +,, 42 and +3 respectively
- 6. In a neutral compound, the sum of oxidation numbers of all the atoms is equili-DEED)
- 7 The oxidation number of an ion consisting of single element is the same a fincharge on the ion e.g. oxidation states of K*, Ca** Ai**, Br*, Si** are +1 +2, +3 1 and 2 respectively
- 8 In ions, the sum of exidation numbers of at the atoms is equal to the charge on the
- 9. In any substance, the more electronegative element has the negative oxidative number

<u>Example !</u>

Determine the exidation number (O.N.) of Mn in KMnO₄ (D.O. Khan Boord, 2007: Hullan Boord, 2010: D.O. Khan Boord, 2011 Rayalp Lahari-Boord, 2013, 2014.

Osidation number of K = +1 Oxidation number of O = -2Let mudation number of Mn = x

$$+1 + x + 4(-2) = 0$$

 $x - 7 = 0$

Thus for KMnO₄

x = +7

College Chemistre: Part-1

Emple 2

Section.



 Determine the oxidation number (O.N) of N in NH₄* Oxidation number of H=+1

Let oxidation number of N = x

$$x + 4(+1) = + 1$$

 $x + 4 = +1$

$$\begin{array}{ccc} x + 4 & = +1 \\ x & = 1 - 4 \end{array}$$

BALANCING OF EQUATIONS BY OXIDATION NUMBER METHOD

This method is based on the principle that

The total number of electrons lost by a reducing agent is equal to the total number of electrons gained by an oxidizing agent.

Following rules are used for balancing equation by orditation number method

- Identify the elements, which undergo a change in oxidation number and write their exidation numbers over the symbols.
- Indicate the number of electrons gained or lost by means of arrows.
- 3. Equate the number of electrons gained and not by multiplying them with suitable numbers.
- Use the suitable numbers obtained in step (III). to become the principle reactable and products.
- Balance the rest of equation by inspection method.

Example,

+ Cu(NO₃₁₈ + NO + H₂O

Identify the elements, which undergo a change in exidation number and write freir oxidation numbers over the symbols

In this equation N is reduced from +5 in HNOs to +2 in NO But it's oradetion state. It may send determine the ... uns equation N is reduced from +5 in HNO $_2$ to +2 in HO out it is obtained in the lit not changed in CurNO $_2$ is. Therefore, write HNO $_3$ twice and determine the humber of electrons gained and lost and equate them.



College Charactery Party

HNO₃ + Cu + HNO₃
$$\xrightarrow{+2}$$
 Cu (NO₃)₈ + NO + H₈O loss of 2 e' × 3 = 6 e'

- Use the numbers obtained in above step to balance Cu and $\mbox{HNO}_{\mbox{\scriptsize 3}},$ Thus, Cu is multiplied by 3 and HNO_4 by 2. $HNO_x + (3 Cu + 2)HNO_x \longrightarrow Cu(NO_x)_x + NO + H_xO$
 - Balance the rest of equation by inspection method. $6HNO_a + 3Cu + 2HNO_a \longrightarrow 3Cu(NO_a)_a + 2NO + 4H_2O$

BALANCING OF EQUATIONS BY IC IOD

It is based on the principle that

The total number of electrons lost by a reducing agent is equal to the total no. of electrons gained by an oxidizing agent.

Following rules are used for balancing equation by ion electron method.

- 1. Write skeleton equation showing only those substances, which are actually involved in the reaction.
- 2. Identify the elements, which undergo oxidation and reduction and split up the reaction into oxidation and reduction half reactions.
- 3. If element undergoing oxidation state is free then it is written as such, otherwise it should be written as motecular or lonic specie
- 4. Balance each half reaction separately.
- 5. Balance oxygen by adding H₂O.
- 6 Balance hydrogen by adding H* ions in case of acidic media and OH ions in case of balance hydrogen by adding H* ions in case of acidic media and OH ions in case of acidic media. basic madia.
- 7 Write down the number of electrons gained and lost in each half reaction.
- Equate the total number of electrons gained and lost by multiplying the two half reactions has electron. reactions by suitable numbers.
- 9. Finally, add the two half reactions to get the resultant balanced equation.
- 10. Simplify the resultant equation if necessary in order to obtain the net conic equation.

college Chamistry: Part-(

Electrocky and

Example I (Acidic media): Cf + MnO_c

-→ CI₂ + M_{D2+} Identify the elements, which undergo oxidation and reduction and split up the D.G. Khan Soned, 2009) reaction into oxidation and reduction half reactions.

550

Balance oxygen by adding H₂O

Balance bydrogen in each half reaction by adding H* low.

Write down the number of electrons gained and lost in each half reaction.

$$8H^{+} + M_{0}O_{4} + E_{6} \longrightarrow M_{0}^{2+} + 4H_{2}O$$
 (1)
 $2CI^{-} \longrightarrow CI_{4} + 2\varepsilon$ (2)

 Equate the total number of electrons gained and lost by multiplying eq.(1 by 2 and eq.(2) b) 5.

$$16H^{+} + 2MnO_{4}^{-} + 10e \longrightarrow 2Mn^{3*} + 8H_{4}O$$

$$10C1^{-} \longrightarrow 5Cl_{4} + 10e$$

Finally, add the two half reactions to get the resultant balanced equation.

To write electrons fort or gained, following formula is used.

Total charge on L.PLS. - Total charge on R.H.S.

If the manner is negative, then the electrons are last. Therefore, add electrons on R.H.S.

If the manner is positive, then the electrons are gained. Therefore, add electrons on L.H.S.

"At in the electrons are gained.

For reduction half reaction, (+8-1)-(+2)=+5, Thus, add 5e on LHS for axidation half reaction, (+8-1)-(+2)=-2, Thus, add 2e on RHS.

College Charactery: Parts!

57U

Example 2 (Basic Media)

$$MrO_4 + C_2O_4^{-2} + H_2O \longrightarrow MrO_2 + CO_2 + OH$$

Identify the elements, which undergo oxidation and reduction and split up the reaction into oxidation and reduction half reactions.

$$M_{II}O_4 \longrightarrow M_{II}O_1$$
 (reduction half reaction)
 $C_2O_4^{2-} \longrightarrow 2CO_2$ (oxidation half reaction)

Balance hydrogen and covygen by adding H_2O and OH^* ions. $2H_2O + M_1O_4^- \longrightarrow M_1O_2 + 4OH^- _____(1)$

$$C_2O_4^2 \longrightarrow 2CO_2$$
 (2)

Write down the number of electrons gained and lost in each half reaction 3e + 2H₂O + MnO₄ - MnO₂+ 4OH ____ (1)

Equate the total number of electrons gained and lost by multiplying eq.(2) by 3 and eq.(1) by 2, and then add the two half reactions.

$$6e^{-} + 4H_2O + 2MnO_4^{-} \longrightarrow 2MnO_2 + 8OH^{-}_{-}$$
 (1)

FLI CHREAL CONDUCTION

Flow of electric current through a medium is called electrical conduction It may be of two types

Or Destrolytic Conduction

The conduction of electricity certified out by ione present in fused or in an aquistitution of my electrolists is collected. solution of an electrolyte is called electrolytic conduction.

e.g. The motion or aqueous solution of NaCl contain Na* and Cl -1 lons. These tolls utilist electricity. conduct electricity.

Mark Characters Part ! 5 Metallia Conduction

The conduction of electricity through a metal due to free al-

All metals are good conductors of electricity

period son

The process is which electricity is used to carry out a non-ap-This process is done in an electrolytic cell.

Domple:

The socium metal is manufactured on industrial scale in Down's cell by the electrolysis of ooten NaCl.

Saazotitaa

The process in which lonic compaunds when fused or dissolved in water split up into larged particles called ions.

NaCl
$$\longrightarrow$$
 Na^{*} + Cl^{*}
PbCl₃ \longrightarrow Pb^{2*} + 2Cl^{*}

GECIRO CHEMICAL CELLS

- These are of two types
- Electrolytic Cellis
- Voltaic or Galvanic Cells

Sectialitie Cells

The call in which electric current is used to carry and a non-spontantial electrolysis call. The provises is called electrolysis.

Timber.

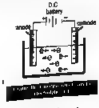
Down's cell. Neison's cell

Continue l'an

I consists of two electrodes dipped into a solution of to electrolyte, containing positive and negative ions. which can move freely in the solution.

Positive electrode is called anode and parative

ties trough is caused supply the called called called called a direct current (D.C.) source, for electrical supply the conflected to a direct current (D.C.) source, for electrical supply the conflected to a direct current (D.C.) source, for electrical supply the called calle







Working

- When electrodes are connected to a D.C. source electric potential is produced in When electrodes are corrected to a survival negative electrode (cathode) and negative potential moves the positive ions towards negative electrode (cathode) and negative or potential moves the positive ions towards negative electrodes (cathode) and negative or potential moves the positive ions towards negative electrodes (cathode) and negative or negative electrodes (cathode) and negative electrodes (cathode) a potential moves the positive ions towards positive electrode (anode). Thus electrolytic conduction occurs in electrols celi
- The movement of ionic charges through the liquid brought by the application of the movement of the charges through the liquid brought by the application of the movement of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the liquid brought by the application of the charges through the charges throug electricity is called electrolytic conduction and the apparatus used is called electroly
- When sons reach the electrodes, electrochemical reactions occur At anode, regnitive sons give up electrons. This process is called oxidation At cathode, positive ions take up electrons. This process is called reduction,
- Hence, an exidation-reduction reaction starts in the cell and vanous products at obtained. The electrons flow through external wire. The electrons flow from another cathode
- The solution will continue conducting electricity until this redox reaction is taking place

The electrochemical reactions that occur at the electrodes during the electrolytic contributions constitute the phenomenon of electrolysis.

For maken slats, products can be predicted. But in ageous solution, if is effort? predict product.

Further electrodes may or may not involve in reaction Following are some products produced from electrolytes

When electrodes are inert (Pt or Graphite).

Unitable	At Controlle	At Associa	
Pb8r _{fm}	Phie	Brees	
NaCl _{et}	None	Class	
NeClas	H _{his}	Claus	
CalCliffer	Cup	Claud	
CuSO	Cupp	Chi	
MeOH _{ingl}	H ₂₄	Cate	
H ₂ SO _{4mg}	H _{au}	Oald	
IQ1O ₂₀₀₄	H _{bul}	Osal	

Continue Paris

When electrodes take part in reaction

(Actiolete disease)	Co Cothage	dealt drawer. Cit dande
CuSO _{Red}	Cu deposits	Cu _{lso} dimotes so forms Cu ²⁺ song
Electrolyte	Ag cathode	Ag saode
AgNO _{sco} and HNO _{sco}	Ag deposits	Aque dissolves to form Aq" yors

INPLANATION OF ELECTROLYSIS

herodesis of Fused solts

- Willien a fused salt of an electrolyte is electrolysed, cations reach the cathode where they take up electrons and become discharged, while
- Anions move towards anode, where they give up electrons and become discharged.

Herhods (Reduction) Pb2+ + 2e- → Pben

<u> Manade (oxidation)</u>

Prend Reaction

$$PbCl_2 \longrightarrow Pb_{M} + Cl_{100}$$

- Similarly fused NeCl and PoBr, are decomposed during electrolysis Fused Pb and Na are collected at cathode and Cless and Brief are collected at anothe respectively
- Electrona flow from anode to cathode.
- Electric current is conducted in the electrolyte by lord.
- Electric current is conducted through the entired circuit by electrons.



Uler troles is al Aquerius Sal area of Solis

When an acueous solution of salt is electrolysed, products are not easily predictable by because H₂O can be both oxidised and reduced and some metal cations are not disc. from their aqueous solution.

Symmole: Electrolysic of Agmeous NaNOs

Consider the electrolysis of an aqueous solution of NaNO.

NaNO_a ionizes as

College Chamistry: Parisi

in MaNO, solution, small amount of hydronium tons (HaO+) and hydroxyl ton (OH+); present due to little ionization of water

i.e.
$$H_aO_{pi} + H_aO_{qi} \Longrightarrow H_aO^* + OH^*$$

Reaction at cathods (Reduction)

When aqueous NaNOs is electrolyzed, Na* ions are not discharged at cathods Install H_aO* ions take up electrons to form hydrogen atoms.

$$H_kO^+ + e^- \longrightarrow H_{kl} + H_kO$$

His atoms then combine to form hydrogen gas.

$$H_{ij} + H_{ijl} \longrightarrow H_{ilije}$$

Thus, H₂ gas is liberated at cathode

Although very less 10⁻⁷ moV dm⁰ of H₂O⁺ ions are present in solution. But as solution H_0O^* is discharged, new H_0O^* ions are produced in solution and thus process continues.

No" lors remain in solution, it is called spectator ion.

Reaction at anode (Oxidation)

At anode, both OH and NOs will be present. However, OH lone are at the discharged

$$OH^- \longrightarrow OH_{(eq)} + \varepsilon^-$$

OH_{eat} then give following reaction

Hence, O₂ is produced at anode. While NO₂⁻ ion is spectator lon.

Order of discharge of lons may also depend upon their concentration.

Overall Reaction

Overall reaction including all the spectator ions.

$$Na^*_{(aq)} + NO_{a^*_{(aq)}} + 2H_{g}O_{g_{1}} \longrightarrow Na^*_{(aq)} + NO_{a^*_{(aq)}} + O_{2(g)} + 2H_{g_{2}(g)}$$

Charles Part

h ciralette Processes of Industrial Important

Exercise Q7

parties the electrolysis of malten sodium chloride, and a concentrated solution of each (Watte Doum's Cell and Nation's Cell)

J. Manufacture of Sodium (Down's Call)

Na is produced in Down's cell commercially by the electrolysis of fixed NaCl between No is produced and graphite anode. Ci_2 is obtained as by product.

MAgade (Oxidation)

$$2Cl^{-}_{01} \longrightarrow Cl_{2G_1} + 2e^{-}$$

A Ganode (Reduction ,

$$2Na^{+}_{0}$$
 + $2\varepsilon \longrightarrow 2Na_{0}$

Na Reaction

I. Manufacture of NaOH (Nelson's Cell etc., Caustic sodia is produced by electrolysis of an eq. solution of NeCl between Titanham annote and Hig or Ag cathode. The process is carried out in Nelson's cell or Castner Kellnercell or Fig - cell

Anode Oxidation.

& Cathode (Reduction)

$$2H_kO_{\mathfrak{g}_k} + 2\mathfrak{g}^* \longrightarrow H_{\mathfrak{s}(\mathfrak{g})} + 2OH_{\mathfrak{g}_k}$$

Net Reaction

Here Cl, and He are obtained as by product.

Na* is not discharged at cathods.

I. Handacture of Mg and Ca by and Ca metals are extracted by the electronals of their fused chlorides. Mg and Ca the collections are extracted by the electronals of their fused chlorides. Mg and Ca

ks collected at cathode and Cl, at anode,

At a extracted by the electrolysis of fused Beuxills (A_kO_k , $2H_kO_l$) in the presence of fused extracted by the electrolysis of fused Hall-Herouli process.

(Na_kAlF_k), This process is called Hall-Herouli process.



5. Manufacture of Anodised Alu Memofrature of Ampensor washing if an anode in an electrolytic cell containing supplies.

Anodized AI is prepared by making if an anode in an electrolytic cell containing supplies. Anodized A) is prepared by meaning of mide is contact over Al. This layer resists compared over Al. This layer resists con The freshly enodized As is hydrated and can absorb dives

6. Reflaing of Copper

Cu in made pure in an electrolytic cell in which impure Cu is made anode and absheet of pure Cu is made Cathode. CuSO4 is used as electrolyte

The atoms of Cu from impure Cu anode are converted to Cut* tons and goes him to solution. From solution Cu²⁺ ions are reduced to Cu metal and deposit on cathode In cathode is made up of pure copper. The impurities are left with anode

7. Electropicting

Copper, Silver. Nickel or Chromism plating is done by various types of electrolytic of In the process, one metal is deposited at the surface of other metal.

Exercise Q13

Describe a galvanic celt explaining the function of electrodes and sait bridge.

VOLTAIC OR GAI VANIC CELL (Zn - Cu Cell)

The cell to which a spontaneous redax reaction is used to produce electric curette nation Voltage or Galacaic cell.

Daniel's cell, Ni-Cd cell, Fuel cell Consider 2n-Cu cali

It consists of two half cells that are electrically connected. In each half cell, half reside takes place. Overall reaction of the call is the sum of two half call reactions.

In each half cell, an electrode is dipped into solution of its ions. Zn electrode is dipped into solution of its ions. Zn electrode is dipped into solution while the indisease in Control is a solution while the indisease in Control is solution. ZnSO, solution while Cu is dipped in CuSO, solution.

Two electrodes are connected by wire, while two solutions are connected by self height Sait bridge consists of an aqueous solution of KCI in gel.

Cu electrode is dipped in 1M CuSO₄ solution in one half cell giving following equilibria:

Zn electrode is dipped in 1M ZnSO₄ solution in other half cell giving juiterium eculfibrium

Two half cells are electrically connected by a set bridge.

When two electrodes are commerced by a wire, electrons start flowing from Zn electrode o Cu electrode.

It is because Zn has creater landered to one electron than Cu. Thus it becomes more negative than Cu

Rescriptor of Zn-Cu sulvanic cell

Atonode (oxidation)

$$Cu_{i}^{R_{0}}_{(aq)} + Zr_{(b)}^{a} \longrightarrow Zr_{(aq)}^{R_{0}} + Cu_{(q)} = 1.1 \text{ V}$$

This Voltair cell can be represented as $Zn_{s0}/Zn^{2+}_{(s=0)}(1M) + (Cu^{2+}_{(s=0)}(1M)/Cn_{(s)}E^{0} = 1.1 \text{ V}$

THE PARTY OF THE P

had but of Salt Realige

ASSET bridge consists of an aqueous solution of KCI in get. It performs following functions

- Itipitings alectrical contact between two half-cells.
- librates to separate the two solutions of half cells. It solutions were missed, direct observed reaction would occur and helf cells would be destroyed.
 - R-maintains electrical neutrality of two solutions.

Duning reaction. In half cell continuously losse sharpers. Thus, in this positive charge is bgeeing while

Cu half cell continuously receive electrons, thus it sees on pollection negative charge.

Oue to collection of positive charge in Zn electrode half cell and collection of negative

Sett bringe prevents the net accumulation of charges in either beater. Thus from charge in Cu half cell would stop the reaction. entire Cu half cell, negative ions diffuse through the salt bridge into the positive Zn half

In voltaic cell the electric current flow through extend circuit. It can be used to hight Thus, it meintains the two solutions electrically neutral tob, drive a motor sto



Culture Chambers Forti-

Revesibility at Gots and at Volume of the

If external circuit is repeated by source of electricity that opposes voltaic on the reaction in voltate call will be reversed and glactrons flow in exposite direction from Cuto's electrode

$$\begin{array}{cccc} \mathbb{Z} n^{1} \circ_{i_{000}} + 2e^{\epsilon} & \longrightarrow & \mathbb{Z} n_{i_{01}} \\ & & & & \mathbb{C} u_{i_{01}} & \longrightarrow & \mathbb{C} u^{10} \circ_{i_{000}} + 2e^{\epsilon} \\ \mathbb{Z} n^{10} \circ_{i_{000}} + \mathbb{C} u_{i_{01}} & \longrightarrow & \mathbb{Z} n_{i_{01}} + \mathbb{C} u^{10} \circ_{i_{000}} \end{array}$$

Now the cell becomes electrolytic cell, because in this, electricity is being used to conout non-spontaneous reaction.

Oxidation occurs at Cu electrode and raduction occurs at Zn electrode.

FIECTRODE POTENTIAL

The potential set up when an electrode is in contact with one malar solution of its set longer 298 K is called Standard Electrode potential or standard reduction patential.

it is denoted by Eº

Examples.

The standard reduction potential of Zn is 0.76V and that of Cu is +0.34V

Explanation .

When a metal electrode is dipped into the solution of its own ion. There may be be sendencies:

Metal atom from electrode leaves the electron on metal and goes into solution. This is exidation process

$$M \longrightarrow M^+ + e^-$$

Metal ion in solution may take up electrons from the metal electrode and deposite atom on electrode. Thus it makes electrode +ve

$$M^+ + e^- \longrightarrow M$$

The -ve or +ve nature of electrode relative to solution depends upon the relative to solution depends upon the tendencies of the two processes.

At last, a dynamic equilibrium is established due to the same rate of two processes. The further potential difference to do the same rate of two processes. no further potential difference is developed.

A Zn ejectrode in a solution containing Zn⁻⁴ ions becomes near A Zn electronic magnitive of the part of the property of the property of the property of the part of t into solution as Zn1+ ions.

The negative charge on the Zn rod will attract an atmosphere of positively charged zinc long around the red to form an electrical double were as shown in the fig.

The equilibrium is represented as

 $Zn_{(a)} = 2n^{2a} + 2a^{-1}$

atmound Guidatton Percental

The potential of an electrode at which aridation takes place under studients is called standard exidation potential

Stepd oil Reduction 15 departed

The potential of an electrode at which reduction taken place under sta editions to called standard reduction potential

it is represented as

E*_= 0.34 volts → Çu Cu** + 2e

For any substance, standard reduction potential is equal to its standard coddation Powerful but with opposite signa

The standard electrode potential of hydrogen is taken as zero.

Position Q10.

Describe the construction and working of standard hydrogen electrods.

Standard Hedrogen Flectrode (SHC)

COMPLINEMON

- * standard hydrogen electrods consists of a glatinum black electrolutically coated with a layer of linely divided relations black.
- The foil: is connected with a platform wire enclosed within a glass. 24.5. Pure 14, gas passes through the case at 1 aim. It is immered in 1M 1400
- n 1M HCl solution
- The Practs as an electrical conductor it also have to stain squilless on the stain squilless of the stain squilles Spalling between gas and its ions in solution.









- Chambers: Part-1

Standard potential of standard hydrogen electrode is arbitrarily taken as sem, be determined by comparing them with sur potentials of all other electrodes can be determined by comparing them with SHE

SHE can act as both cathode and anode depending upon other electrode with which to connected.

Pollowing are the reactions of SHE

SHE on Anodes

$$2H^* + 2e^*E^*_{ee} = 0.00 \text{ volta}$$

SHE or Cathode:

$$2H^*+2e^*\to\ H_x$$

$$E^0 = 0.00$$
 volts

Measurement at the ende Perential

Unionunetely. It is not possible to determine the single electrode potential

However, the potential of a single electrode can be determined by coupling it with standard electrode of known potential such as standard hydrogen electrode (SHE) in I galvanic cell. The two solutions are exparated by a porous partition or a sak bady containing conc. solution of KCI. Salt bridge provides a highly conducting path between its two electrolytic solutions. The potential difference is measured by a voltmeter

Since standard electrode potentials (oxidation and reduction) of SHE are taken at ten Therefore, the measured value of cell potential will be the potential of unknown electrods

In this way, a series of standard reduction potential can be built up for different electrodes.

Csample 1: Measurement of Zu Electrode Potential

Electrode potential of Zn electrode can be measured by making a galvanic cell it ones of Zn electrode dipped in 1M solution of its ions and SHE at 25°C. Under the studied conditions unless the studied of the studied conditions. conditions, volumeter shows 0.76 volts and the deflection shows that electrons are flowed

Thus, Zn has greater ability to give up electron than He gat by 0.76 volts.

The reactions are

At Anode (oxidation)

Since Standard electrode potential of SHF is 0, therefore 0.76 is the standard or being length of Sh. potential of Zn.

Hence, its standard reduction potential will be -0.76 yells.

sample 2. Measurement of Car Hychode Potential Electrodic potential of Cu electrode can be measured by making a galvanic cell. It consists of Cu electrode is dipped in 1M solution of its ions and SHE at 25°C. Under the standard conditions, volumeter shows 0.34 voits and the deflection shows that electrons are flowing from SHE to Cu.

5)*

Thus, H_{g} has greater ability to give up electron than C_{d} gan by 0.34 volu.

The reactions are

At Anode (oxidation)

At Cathode (reduction)

Stace Standard electrode potential of SHE is 0, therefore 0.34 is the standard reduction potential of Cu.

Hence, its standard existation potential will be -0.34 volts.

OR LITETROCHEMICAL SERIES

When electrodes are arranged in order of their standard electronic spirogen souls, the resulting list is called Electrochemical Spries.

According to International Union of Pars and Applied Chemistry (IUPAC), standard decirode potential are given in reduction mode.

The conduction potential of the electrodes can be obtained by revening the sign of their reduction potential.

These reduction potential are given under standard conditions. i.e. 25°C and 1 sen were and 1 M solution of its ions. Changes in temperature, concentration and pressure will change the values of reduction potential.

A brief sketch of this series is given below





Table (10-2), standard is ductioned stendardy. First element at 298K.						
l lemen‡		t he tree	lı-	ja – Egorapia	Standard reduction potential (1-)	
Li	d*+ e−	→	u		-3.045	
ĸ	K*+ e	-	ĸ		-2.925	
Q	Qa ^{k+} + 2a ⁻	\rightarrow	Ca.		-2.87	
Nu	Nn"+ e"	\rightarrow	Na		-2.714	
Mg gr 1	Mg ⁴ *+ 2#*	\rightarrow	Mg		-2.37	
A	Al ⁶ * + 8e	\rightarrow	Al ,	ts	1.66	
Za g	2n*+ 24**	\rightarrow	Ze	2	-0.76	
C- 10	Qr5+ 3e-	\rightarrow	Cr) pg	-0.74	
Pa B	Fe ²⁺ + 2e ⁻	•	Fe	8	-0.44	
C4 ₽	C4*+ 2**	\rightarrow	Cd	표	-0.408	
N	Nf ² *+ 2e ⁻	\rightarrow	N		-0.25	
Sa 🖥	Sa*++ 2a**	*	5n	sa ilignatia grianaria	-0.14	
Pb 2 *	Pb*+ 24**	\rightarrow	Pb	1	-0.126	
Ha	2H*+ 2#		, H ²		0.000	
O ₄	Ox** + 2a**		Сę		+0.34	
Çn	ርህነት ፈግ	\rightarrow	OL.		+0,581	
1,	$l_4+2\pi^-$	\rightarrow	21-		+0,535	
Fe	Fe ^{lly} + 3e	\rightarrow	Fe		+0.771	
Ag	Ag*+ e™	\rightarrow	Ag		+0.7994	
Hg	$Fig^{0a} + 2e^{-}$	->	Hg		+0.885	
Brg	Be _p + 2e ⁻¹		28c	-	4-1.08	
Cla	O ₀ + 2a	-	201	-	+1.360	
Au	Ad ³⁴ + 341	-	Au		+1.50	
Fit	F2+ 24"	\rightarrow	žP-		+2.87	

Order Chambion Restal

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PPLICATIONS OF FITCHFORD MEATSURES

The emil series has many applications.

1. Conferming the leavibility of a the moral Reaction

Using standard reduction and exidation potentials, spontaneity of a self reaction can be determined.

The call voltage is determined using the formula

$$E^0_{-} = E^0_{-} + E^0_{-}$$

- E°_m = E°_m + E°_m0

 if E°_m is positive, cell reaction is strontaneous.
- . If E all is negative, cell reaction is non-spontaneous.

Example-1

$$Z_{\Gamma} \longrightarrow Z_{\Gamma}^{0+} + 2e^{-}$$
 $E^{0}_{0+0} = 0.76$ volts $E^{0}_{0+0} = 0.34$ volts $E^{0}_{0+0} = 0.34$ volts

$$Z_n + Cu^{2+} \longrightarrow Z_n^{2+} + Cu$$
 $E^1_{ad} = 1.10 \text{ volts}$

This reaction is apontaneous. However, its reverse reaction will not be spontaneous.

Thus, Cue+ can exidize solid Zn, but Zn*+ cannot exidize solid Cu.

2. Colculation of Cell Victorio (end)

While making Gaivards cell, the electrode above in electrotherical series will act as along, while the electrode below will act as cathode.

At anode exidation takes place, at cathode reduction takes place. The cell potential is determined by the formula

Winfor Zn.- Cu cell

Origination potential of Zn is ± 0.76 volts, while reduction potential of Or^{10} is 0.34 volts, thus

$$E^{o}_{cold} = E^{o}_{(col)} + E^{o}_{cold} = 0.76 \pm 0.34 = 1.10 \text{ volts}$$

Electromotice Force (em) The sail voilings or amf resources the jove with which steeders more in the entered rate.

Thus, it measures the tendency of the cell reaction to takes place. Hence, Galvanic cell segments of the reactions to occur. thus, it measures the tendency of the cell reaction to become sheet quantitative measure of the relative tendency of the reactions to occur.

Example: The end of a Zn-Cu call 5 1 10 volts



Colon Charleto: Patel

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3. Comparison of Relative Temperary of Metals and Non-Metals to Get Oxidison of Reducat

Greater the value of reduction potential of a speale.

Greater is its tendency to accept electron.

Hence it will be a better oudding agent

Ехаптова

ions like Au^{a+}, Pr^{a+}, Hg²⁺, Ag⁺ , Cu²⁺ and non-metals like F₂ , Cl₂ , Br₂ and I₂ which <u>is</u> below SHE have more lendency to accept electrons. Hence, these easily undergo reduction. end series show

Strong oxiditing accents have lause positive values of standard reduction potential. These he below SHE in the series

While

Strong reduction poents have large penaltive values of standard reduction potential There he above SHE in the series.

4. A fata e Chemical Remainite of Metals

Greater the value of standard reduction potential, leaser is the ability to loose electron (i.e. to change into positive lons). Hence its reactivity is less, Thus

- Metals tike I.I., Na., K and Rb are highly reactive.
- Coinage metals Cu. Ag and Au are the least reactive because they have positive seduction potential.
- Metals like Pb, Sn, Ni, Co and Cd which are very close to SHE react very slowly with steam to liberate hydrogen gas.
- Metals the Fe, Cr, Zn. Mn, Al and Mg have more negative reduction potential. This these react with steam to produce metallic oxides and He gas.

5. O action of Metals with Dilute Acid

Greater value of reduction potential. Letter is the ability to loose electron to just positive ion. Hence weaker is its tendency to displace Ha.

Thus any metal pixove SHE could displace He from acid but those below it san its place He.

Metale like Zn, Mg and Cu have negative values of reduction potentials. Thus, these call liberate He liberate H₂

Metals like Au. Pb. Ag and Cu have high positive values of reduction potentials. This these do not liberate H_0

Chamistry: Port-1

6. Displacement of One Head by Swother lead as Substant

Any metal can displace any other metal from its solution, present below in the electrochemical series.

Thus, Fe can allaplace Ou from CuSO.

Fe + CuSO₄ --- Cu³⁺ + FeSO₄

However, Zn cannot displace Mg from MgSO.

Zn + MgSO_e → No Reaction

MODERN BATTLEIES AND FUEL CILLS

These may be of two types

Promate creas

These can not be re-charged.

e.g., Alkaline battery, Silver-oxide battery

\$110M0ARFC11[8

These can be re-charged

e.g. Lead accumulator

Examples:

battery.

Alkaline battery, Stiver-oxide

Difference between Primary and Secondary Colin.

		Secreptive Lefts
Primmt Colls		They can be recharged
They cannot be recharged	-	The second state of the second state of the second
They contain electrolyte in an	1	
adsorbent or separator. Thus these		are the same property of Californ (1991 Californ).
are called drt. cells.	-	They are generally larger in size.
They are generally smaller in size		
	- [They require regular maintenance and
They are disposable		ratherging

Examples: end Accumu



College Chembers: Port-L

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Electrodon by his

LEAD ACCUMULATOR ALCHARGI ABLL

It is a re-chargeable car battery

Construction

A fully charged battery consists of A cathode of lead oxide (PbOL) And Anode of Metallic Lead

The electrolyte is 30% H-SO, with density 1.25 d/cm2.

d gride I Bed (all not of visit saper. Lead one to reco-

Two electrodes are connected through an edemai circuit. It produces electrical current by discharging.

-0.00

It produces electric current by discharging

Discharging

At Anode (oxidation)

At anode, lead atoms loose electrons and orddise to Phi³⁵ Phi³⁴ Jops combine with 50.2 ions and deposit as Pb5O4 on anode.

The electrons released pass through an external chault and thus produce electricity

PbOs receive electrons from anode. It then gives region reaction with H° tors in the At Cathode freduction). electrolyte to give lead ions and water

The Pb⁴⁺ ions then combine with the SO₄⁴ and deposit as PbSO₄ on cathode

The overall reaction is

Thus, electron flows through external circuit and can be used to light bulb etc. Admit produce about 2 volts. cell produce about 2 volts.

When both electrodes are completely covered by PoSQ, reaction stops and III. III. current is produced. The battery is then recharged

Office Chambers: Part-I

Re-alexystres

During this, anode of the external electrical source is connected to anode of cell while cathode is connected with cathode of the cell.

Thus, redox reactions are reversed.

At Anode (This becomes cathode and reduction occurs during re-charging)

→ Pb₍₄₎ + SO₄² ₍₄₄₎ PhSO_{Me} + 2 g

<u>ArGothode (This becomes anode and axidation accurs during re-charatral</u>

Overall Renation

$$2P_0SO_{4\mu_0} + 2H_1O_{70} \longrightarrow P_0G_0 + P_0O_{1\mu_0} + 4H^{+}_{[\mu_0]} + 2SO_4^{-1}_{[\mu_0]}$$

During discharging, the cons. of acid falls and its density decreases to 1.15 g/cm³. During recharging concentration of acid goes back to 1.25 g/cm³ and voltage of call is come back to award 2 volts.

 $\widehat{\mathbb{H}}$ is matrily used as car limitary for starting angine, for lightening up car lights etc.

PSPICAL CAR BATTERY

- A typical car battery consets of six calls, connected in series
- Each cell gives 2V.
- Each cell contains two lead gifts packed with the electrode materials.
- This anode is spongy isad and cathode is countered PbOs
- The grid is immersed in an electrolytic solution of nearly 4.5 M H₂SO₄ (30%) Fibre Sizes sheets between the grids present shorting by addicted physical contact
- When the cell is discharged, it generates electric energy as voltace cell.

A RALINE BATTERY INON-RECHARGEABLE

Constituentian

itis a.dry alkaline cell. bittes Zn and MnO, as reactants.

Za rod acts as anode, while

MnO₄ acts as cathode

Electrolule consists of KOH southorn

Therefore, it is alkaling.



The bettery is enclosed in a gire; contacter.

In a passery is empower as a mass to give larger effective area, it allows cell to deliver more current than common dry cell. It also has longer life.

A saking and Heartons

its reactions are

At Anode condation)

At cathode (reduction)

The overall reaction is

$$Z_{\Pi(S)} + 2M_{\Pi}O_{S(s)} + H_{2}O_{(s)} \longrightarrow Mn_{2}O_{S(s)} + Zn(C(H)_{S(s)}$$

Cell voltage is 1.5 V

SHALP OXIDE BATHERY

It is used as power source in electronic devices e.g. watches, auto exposure cameras and electronic calculators. It is very small in size.

Constitution

The cathode is of silver oxide Ac.O The anode is of Zn metal.

The electrolyte is basic.

Following reactions occur

At anode (oxidation)

$$2n_{sa} + 2OH_{sap} \longrightarrow 2n(OH)_{2sp} + 2a^-$$

At cathode (reduction)

$$Ag_2O_{eq} + H_2O_{gg} + 2e^- \longrightarrow 2Ag_{bq} + 2OH^-_{(eq)}$$

The overall reaction is

$$Ag_2O_{int} + H_0O_{int} + Zn(s) \longrightarrow 2Ag_{int} + Zn(OH)_{Bith}$$

Voltage of battery is 1.5 V

Charles Park

CIBIL CADMIUM CELL IRI CHARGI ABILI Ris a strong cell largely used in NICAD or Nicke Cadmium battery

The anode is of Cadmium.

The cathode is of NiO.

The electrotyte is alkaline.

furling and Reactions

Following teactions occur

MAnode foxidation.

Cd_(e) + 2OH^{*}_{[ea)} ---> Cd(OH)_{[eq)} + 2e⁻

Monthode (reduction)

$$N|O_{2aa} + 2H_2O_{(l)} + 2e^- \longrightarrow N_l(OH)_{2aa} + 2OH_{lea}$$

lle Resellon

$$Cd_{od} + NiO_{2|o\rangle} + 2H_{z}O_{(b)} \longrightarrow Cd(OH)_{0|o\rangle} + Ni(OH)_{3|o\rangle}$$

<u>Voltage</u> of the cell is about <u>1.4 ∨</u>

in this, solid reaction products are deposited on electrodes like read bettery. Thus, mation is easily reversed during recharging. Since no sales are produced during charging their story, therefore, it can be sealed it is light weight.

- It is used in battery operated tools and portable computers.
- it is also used in cordless razors, photoflash units

WI CLIAS

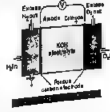
These cells, fuel gases H_{z} and O_{z} are reacted and estricity is obtained.

within (ion

Destrodes consists of hollow tubes made of porous interest carbon coated with Pt.

Placts as cotalusi.

Electrolyte is KOH.



area (in In a common fixed call, H_0 is oxidized to H_0O and O_2 is reduced to OH^* tons.



Charleter Paris

At Anode (oxidetion)

$$H_{\text{tim}} + 2OH_{\text{tim}} \longrightarrow 2H_{\text{g}}O_{\text{m}} + 2e^{-}| \times 2$$

At Cathode (reduction)

Overall reaction

- These cells run continuously as long as reactants are supplied.
- These cells are operated at high temperature
- These are used in space vehicles. Water is formed as by product. It may be content and used for drawing. Thus Firel cells produce electricity and pure water during sec flights.
- To get asveral Kilowatts of power, many fuel cells are connected together
- Fuel cells are very efficient. They convert about 75% of fuels bond energy into exting
- Fuel cells are light, portable and acurces of electricity.
- Many fuel cells do not produce polkrinnts.
- Some other cell reactions in fuel cells are
 - \longrightarrow N₂ + 3H₂O 1 2264 + 3/2 On
 - → N₂ + 2H₂O
 - 2 NH + O2 ---> CO₂+2H₂O 3. CH₄ + 20₂

Chapter Park OBJECTIVE AND SHORT ASSILTE OFF SHOPS IT OTOSSE

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Multiple choice questions.

The outhodic reaction in the electrolysis of dil. H_2SO_4 with Pt electrode is

(a) reduction (b) oxidation

- (a) rectionance (d) Neither exidation nor reduction (d) Neither exidation nor reduction (e) Bord, 2009 (Februahad Board, 2009) (Segodia Board, 2017, (2).6. (five Board, 2012) (C.)
- (ii) Which of the following statement is correct about galvanic cell.
 - (a) anode is negatively charged (b) reduction occurs at anode
- (c) cathode is positively charged (d)reduction occurs at eathode (s.G. Ron Board, 2012) (Gargodia Board, 2012) (Leitora Board, 2013)
- (a) Stronger the caldleing agent greater is the

(b) reduction potential (a) oxidation potential

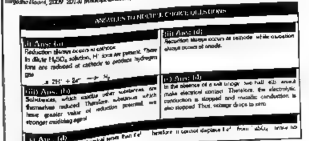
(d) EMF of cell u and 2004)

- (c) redox potential (d) EMF (dual Bound, 2010; Gubernook, Board, 2013, 2914) (Sugar
- (b) If the oult bridge is not used between two half calls, then the voltage (b) decrease slowly
- (c) does not change (d) drops to 2010 (c) does not change (d) drops to 2010 Finished Board, 2007) (Rahasalper Board, 2009) (D.C. Aften Board, 2009) (D.C. Man Board, 2012) (Rahasalper Board, 2012) (Sagodha Board, 2012) (E) Briddon Board, 2011 2012) (Sargodha Board, 2012)

- if a strip of Cu metal is placed in a solution of FaSO, (b) Fe is precipitated out

(a) Cu will be precipitated out

(c) Ou and Fe both dissalve for 2009 2013 planeted abort 2019 (Out and Fe both dissalve for 2019) (Outramed Boort, 2019)



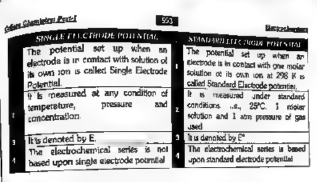
Fit1 in the blacks with suitable words. In OF2 and is ___ in H_2O_3 . The axidation number of O atom is ___ in O_3 . Q2.

592 College Chantery, Paris (ii) Conductivity of metallic conductors is due to the flow of Reaction taking place at the ____ is termed as oxidation, and at the electrolytes is due to flow of called reduction s setup when a metal is dipped in its own sons the Cu-cathode when electrolysis is performed for 0.50, fast. (v) Cu metau solution with Cu-cathodes. ____ volts end its oxidation potentia; (w) The reduction potential of Zn is ____ __ volts. react together in the presence of In a fuel cell. Answers: (ii) electrons tens (iii) anode cathods (iii) +2. 1
(b) equilibrium (c) deposits on (c) H, and O, PI ratalyst Q3. Indicate TRUE or FALSE on the case may be. In electrolytic conduction electrons flow through the electrolyte. In the process of electrolysis, the electrons in the external circuit flow from calificaä Sugar is a non-electrolyte to solid form and when dissolved in water will also be passage of an electric current. A metal will only allow the passage of an electric current when it is in cold state The electrolytic products of aqueous copper (II) chloride solution are copper dilogram Zinc can displace son form its solution. SHE acts as cathode when connected with Cu electrode viii) A voltaic cell produces electrical energy at the expense of chemical energy Lead storage battery is not reversible cell. Or changes in oxidetion number when K2Cr2O₇ is repoled with HCL in) False (iii) False (iii) False (iii) True Q7.Describe the electrolysis of molten sodium chloride, and a c

Q0.80 hat to the difference between single electrode potential and standard of

potential? How can it be measured? Gloc its importance.

sodium chloride Solved on Page 575



95.Dualine the Important applications of electrolysis. Also write the electrochemical reactions impointed therein. Discuss the electrobets of CuSO, using Cu electro-AgNO solutions using Ag electrods.

910. Describe the construction and working of standard hydroges electrode. Solved on Page 579

QII Let the population $Fe^{g+}+Ag o Fe^{g+}+Ag'$ appulational \emptyset not, write appelaneous

reaction involving these species. Selection Fe³⁺ In this reaction, Fe is reduced white Ag is caldized. Therefore, Fe^b well act as

cathode while Age as anode Thus, emit of the cell will be

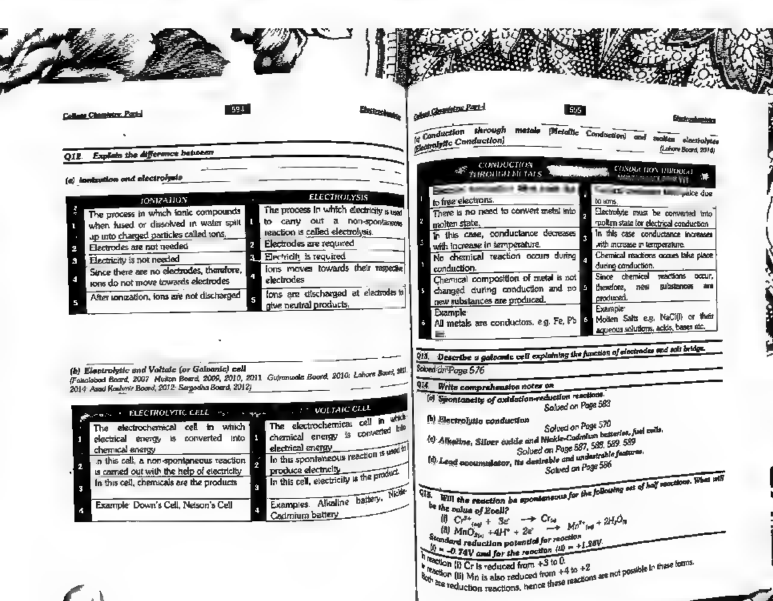
a

$$E^*_{max} = F^*_{max} + E^*_{max}$$
 $F^*_{max} = -0.7994 + (-0.44)$
 $E^*_{max} = -0.7994 - 0.44$

E' = - 1.2394

Since earl of cell is negative, therefore the cell-reaction is non-spontaneous But if the electrodes are reversed, the cell-reaction becomes spontaneous as

Por this E* = + 1 2394



-01



Fist

However, if reaction (i) is revenued so that Cr is oxidized then the reaction to

emf of the cell is given by

the cell is given by
$$E^*_{out} = E^*_{out} + E^*_{out}$$

$$E^*_{out} = +0.74 + 1.28$$

$$E^*_{out} = | x.02 \text{ V}_{out}|$$

Q16 Explain the following with receases

(a) A poroue plate or a soft bridge is not required in fined scorage call. fl.ahore Board, 2009 2010. Sargodha Board. 2007 2009. 2014. Fetsolabod Board, 2007, 2011 Bale Board, 2011 D.C. Khen Board, 2012 Reuvalpinel Board, 2012 Multan Board, 2012)

A sall bridge has two main functions

- It joins the solutions of two half cells and thus complete the circuit
- It maintains electrical neutrality of the two half cells as ions can pass through it.

In Lead storage battery, both cathode and smode are dipped in the same Kilder Therefore, excess positive or regative ions are not produced in the solution. Hence that no need of salt bridge.

(b) The standard axidation potential of In is 6.76 V and its reduction potential is 4.5 (Lahore Board, 2008, D.G. Khen Board, 2019)

According to the law of conservation of energy, energy can neither be created to destroyed Therefore, if standard exidation potential of Zn is 0.76 V, then its potential of Zn is 0.76 V, then its potential of Zn is 0.76 V. reverse process, i.e. standard reduction potential will also be same but with positive of Thus

(Remaipindi Boerd, 2007: Guranaski Board, 2009: Bahawalpur Board, 2009: Palaolabad Board, 2011: Pil Khan Board, 2011: Asad Kashmir Board, 2012: Sargodha Board, 2012: 2014; Zn can displace hudroasen from addition

Zn can displace hydrogen from acids but Cu cannot die sa. Why?(automic lie gamentale Report 3013)

No can displace hydrogen from acids but Cu cannot. Why? gautee Boord, 2011.

Greater the value of reduction potential, Lesser is the ability to loose electron to be strong the strong to the control of the strong to the strong to loose electron to be strong to the strong to loose electron to be strong to the strong to loose electron to be strong to the strong to loose electron to be strong to the strong to loose electron to be strong to the positive ion, Hence weaker is its tendency to displace Ha-

Callent Chemisters Parts

1947

- Metals like Pt, Pb, and Cu have high positive value of reduction potential. Thus these do not liberate Ha
- . Metals alice No and K have negative values of reduction potential. Thus, these can Iberate Ha.

(d) The equilibrium is see up between main atoms of eleginode and tone of main in a cell.

When a metal electrode is dipped in the solution of its own tons. There may be two undencles

Mates atoms from electrode leaves the electron on metal and goes into solution. This is addation process

Metal ions in solution may take up electrons from the mates electrode and deposit as atoms on electrode. This is reduction process.

At last, a dynamic equilibrium is established due to same rate of two processes. The potential set up when an electrode is in contact with solution of its own for is called Electrode Potentia:

(a) A soit bridge matrixing the electrical neutrality in the cell. Fulnalahad Scorel, 2010: D.G. Khar Seard, 2016: Onl-monte Scorel, 2010: In Scorel, 2013: Aund Kanbush Bourd, 2013: Martin Scorel, 2029, 16139

Two half cells are electrically connected by a sait bridge.

During reactions of this cell, In hair cell continuously toose electrons. Thus, is this positive

While, Cu half call continuously receive electrons, thus it goes on collecting negative charge.

Collection of positive charge in 2n electrode half cell and collection of negative charge in

Sait bridge prevents the net accumulation of charges in either basise. Thus from harding prevents the net accumulation of charges in either basise. Thus from harding into the positive Zn half call negative Ca half only negative hardings the two solutions, electrically neutral.

(i) Lead accumulator is a charge ship bettery. W Leard accumulation is a survey of the control of

mesualphind Bodrd, 2010: Mustin source areas source source, 2013: Sorgoon Heerd, 201 Off Expirite through equations how issed battery is recharged? (Lehem Board, 2014)



College Chapters Pro-I

For recharging of Lead Accumulator, the anode of the external electrical source connected to snode of cell while cathode is connected with cathode of the cell.

Thus, redox reactions are reversed

At Anode (This becomes cythode and reduction occurs during re-charaing)

$$PbSO_{4a4} + 2 \pi$$
 \longrightarrow $Pb_{64} + SO_4^{B}$ $_{beg}$

At Cathods (This becomes anode and ordention occurs during re-charginal

Overall Renation

$$2PbSO_{440} + 2H_2O_{60} \longrightarrow Pb_{44} + PbO_{844} + 4H^*_{440} + 2SO_4^{2}_{(44)}$$

During discharging, the cond. of acid falls and its density decreases to 1.15 g/cm² During re-charging concentration of acid goes back to 1.25 g/cm² and voltage of call is come tasks

(a) Impure Cu can be purified by electrolytic process.

(Cityrenwede Beard, 2009; D.G. Khen Beerd, 2009; Lahore Beard, 2012; Bahawalpur Board, 2012; Fainktel Board, 2013; Cityrenwola Board, 2014)

Impure Cu can be made pure in an electrolytic cell. Thick sheets of impure copperate made smode, while, thin sheets of pure copper are made cathode in the cell. These sheets at placed in an electrolytic solution of CuSO4.

When current is passed through the cell, Cu from anode is exidized to Cu²⁺ ions, which go mito the solution. From the solution, Cu²* ions are reduced to metallic Cu and deposits # pure Cu on cathode. In this way, impure sheets of Cu (anoda) become thin, while pur sheets of pure Cu (cathode) become thick

The reactions in the cell are

At Anode (oxidation)

V COOMES

Thus, there is no net reaction in the cell. However, the net result is the purification of Qu OH S.H.E. cote so anode when connected with the Cu electrode but as enthals with the

[Multen Board, 2008, 2011 Sargocha Board, 2016]

- 5 H.E. (0.0 voits) has higher reduction potential then 2n (- 0.76 voits). Thus, when both these are connected electrons and and these are connected electron flows from Zn to S.H.E. Hence, Zn acts as another set.
- 2 S.H.E. (0.0 volts) has lower reduction potential then Cu (+ 0.34 volts). Thus, when lower these are connected electron flows from S. H. E. H. E. H. Cu (+ 0.34 volts). Thus, when lower the same statement of the same statement of the same statement. these are connected electron flows from S.H.E. to Cu. Hence, S.H.E. acts as an of Cu. as cathode.

Chamber: Part-I

551

NUMERICAL PROBLEMS & wreset Q & (c) Calculate the axidation number of Chromium in the following.

Thus

CO,

CyCly (Lobors Board, 2010; Multan Board, 2015) K,CcO, Oxidation number of K =+1

Oxidation number of Cl = -1Oxidation number of Cr = xOxidation number of Cr.

can be calculated as For CrCle x + 3(-1) = 0

x - 3 = 0or x = +3

 $K_{\mu}C_{T\mu}O_{\gamma}$

Oxidation number of K = +1 Oxidation number of 0 = -2Oxidation ourober of Cr= x

Oxidation number of Crox x

Thus Por Cr₁O₂

2x + 3(-2) = 0

 $x=\pm 6/2=\pm 3$

2x 6 = 0

Oxidation number of Q = -2

Oxidation number of Cr = x

x - 6 = 0

Oxidation number of O = -2

Oxidation matcher of Cr = x

= +6

For $K_{\mathbf{c}}C_{\mathbf{c}}O_{\mathbf{c}}$ 2(+1) + x + 4(-2) = 0

Fox Cr₂O₇ *

Fox Cr₃-L_y

2x + 7(-2) = -2

2x - 14x - 2 + 14

ox x = +12/2 = +6

Rabor Boord 200: Multim Boord, 2019 (Faint)

300: Robinspins Boord, 2018: capture Boord, 2013)

Ct₂(\$0_{0.2} Oxidation number of \$ = +6 Oxidation number of C=2Oxidetion number of Cre x For $Cr_2(SO_2)$, 2x + 3r(+6) + 4r - 2r = 0 2x - 6 = 0

or x = 6/2 = +3 or Berry 20/13

Thus



2.00

College Characters, Bartel (d) Calculate the catheties numbers of the elements underlined in the join Oxidation number of Na. + 1 Ne.CO, Oxidation number of $O = -2^{-1}$ Oxidation number of O = -2Oxidation number of P = xOxidation number of Nam +1 Oxidation number of C = x Thus For Na₂CO₃ For Na PO. 2(+1)+x+3(-2)=03(+1) + x + 4(-2) = 0x - 4 = 0x - 5 = 0or x = +4or x = +5 $Ca(CIO_3)_2$ (Motion Board, 2006) Californ number of $Ca=\pm 2$ Cr₂(**gO**₄)₂ Oxidation number of Cr=+3 Oxidation number of O = -2 Oudation number of O = -2Oxidation number of Cl = x

Oxidation mamber of S = xThus For Cra(SO₄)₀ 2(+3) + 3((x) + 4(-2)) = 03x - 18 = 0or x = 18/3 = +6

That For Ca(CIO₂)₂ [+2]+2[x+3[-2)]=0[2+2x-12=0]x = 10/2 = +510 FileO₃ (Fatrolphod Board, 2011)

Kalino, Oxidation number of K =+1 Oxidation number of O = -2Oxidation number of Mn= x Thus For KyMnO₄

2(+1) + x + 4(-2) = 0x - 6 = 0or x = +6 For HNO (+1)+x+3[-2]=0x - 5 = 0x w +5

(Resolvinch Search, 2010) Oxidation number of H =+1 Oxidetion number of O = -2Oxidation number of P= x Thus

For HPO₈ (+1)+x+3(-2)=0x - 5 = 0OK X = +5

Oxidation number of H=+1 Oxidation number of O = -2Oxidation number of N= x Thus

> Try Yourself Han Caroch Halos Za(OH), H.PO. Neck

College Chamber Parts

50-

95 (b) Balance the following equations by orderine member method

 $HINO_1 + HI \rightarrow NO + J_1 + H_2O$ Identify the elements, which undergo a change in ordination number and write the midation numbers over the symbols

114 2140 11-1 HINO + HI -- NO + I + H₂O

Determine the no. of electrons gained and lost and equate them.

gain of 3 et x 1=3 et (reduction) HINO, + HI - NO + 1 + HO loss of 1 et ± 3 = 3 et (explanant)

Balance loss and gain of electrons by multiplying HI by 3
HINOs + 3HI → NO + (s + H₂O)

Balance the test of equation by impedian method 2HNO₆+ 6HI -->2NO + 3I₆+4H₂O

Br₄ + NaOH - NaBrO₂ + NaBr + H₂O (Sequence Second, 2018)

Identify the elements, which undergo a change in culdation number and write sixth ordidation numbers over the symbols

0 Br_a + NaOH ---> NaBrO₆ + NaBr + H₆O

Since Br₂ is involved both in oxidation and reduction, therefore, we shall write the Br— butce. Then determine the not of electrons gained and fost and equals them.

Brg + Brg + NaOH -+ NaBrOa + NaBr + HaO ton of 2(b) of a 1 a 10 of (axid

ORIG + BIG + JERBUH - 2NABRO, + 10NABR + 6H₂O
Or 6Bra + 12NACH - 2NABRO, + 10NABR + 6H₂O
Or 35ra + 6NACH - NABRO, + 5NABR + 3H.1 NaBrOs + 5NaBr + 3H₈O

College Chambridge Part (

14 to

Zn + HNO₃ --- Zn(NO₃)₀ + NO + H₂O (NOTE: Continued Report, 2011; Maltin Based, 8018; Margatha Based, 8011; (NOTE: Or may be present in place of \mathcal{Z}_{ab}

identify the elements, which undergo a change in oxidation number and write the caldation numbers over the symbols.

In this eq. N is reduced from +5 in HNOs to +2 in NO. But it's oxidation state is not changed in Cu(NO_a)₂. Therefore, write HNO_a twice and determine the number of electrons gained and lost and equale them.

gain of 3 er × 2 = 6 er (raduction)

$$\downarrow \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \qquad \downarrow \qquad \qquad \qquad \qquad \qquad \qquad \downarrow \qquad \qquad$$

Like the multiplier obtained above to belance loss and gain of electrons. Thus Cu is multiplied by 3 and HNOs by 2.

(Surpodhe Souré, 2012) MnCig + HGi MnCig + HyO + Cig lidentify the elements, which undergo a change in exidation number and write that ordidation numbers over the symbols

In the as, C) is reduced from -1 to MCI to 0 in CI, But it's origination state is not changes in MinCig. Therefore, were MCI torce and determine the no-of electrons gand. and user and abuse from

se the emphysical extraordical alternation of managed game of infactional

College Chambelow Part-1

Balance the rest of equation by inspection method 2 HCl + MnO₂ + 2HCl → MnC₂ + 2H₀O + C₄

$$FcSO_4+K_4Cr_4O_3+H_4SO_4 \longrightarrow Fc_6|SO_{1/3}+Cc_6|SO_{1/3}+K_4SO_4+K_6O_5$$

 Identify the elements, which undergo a change in oxidetion number and write their exidation numbers over the symbols

$$\stackrel{*}{\text{FeSO}}_4 + K_z C r_z O_y + H_0 S O_0 \ \longrightarrow \ \stackrel{*}{\text{Fe}}_{S/S} (S O_4)_s + C r_z |S O_4)_s + K_0 S O_4 + H_4 O_4$$

Determine the no. of electrons gained and lost and equate them

$$\label{eq:control_ent$$

Use the multiplier obtained above to balance loss and gain of electrons $6FeSO_4 + K_aCe_2O_7 + H_aSO_4 \longrightarrow 3Fe_1SO_4)_a + Ce_1SO_4)_a + K_aSO_4 + H_aO_4$

Balance the rest of equation by respection method. $6FeSO_4 + K_eCr_gO_y + 7H_gSO_4 \longrightarrow 3Fe_gISO_{41} + Cr_1ISO_4)_1 + K_gSO_4 + 7H_gSO_4$

Identify the elements, which undergo a charge is ordation number and write time oxidation numbers over the symbols.

In this equation S is radical from $+\delta$ is H_0SC_0 to +4 in SC_0 But it's ordered state is not determine the no. of elections of the shape of the state Offices and lost and equate them.

$$H_2SO_4 + Cu + H_2SO_4 \rightarrow CuSO_4 + SO_7 + H_4O$$

$$loss 4/2 = \pi/1 + 2\pi touldhord$$

Coffee Chambers Paris

- Use the multiplier obtained above to balance loss and gain of electrons → CuSO₄ + SO₂ + H₂O H₂SO₄ + C₀ + H₂SO₄
- Belence the test of equation by inspection method. -> CuSO₄ + SO₂ + 2H₂O H₂SO₄ + Cu + H₂SO₄

Or
$$O_0 + 2H_0SO_0 \longrightarrow O_0SO_0 + SO_0 + 2H_0O$$

 $H_2SO_4 + HI \implies SO_2 + I_2 + H_4O$ identify the elements, which undergo a change in oxidation number and write their oxidation numbers over the symbols

$$H_2SO_4 + HI \longrightarrow SO_2 + I_2 + H_1O$$

Determine the no. of electrons gained and lost and equate them

$$\begin{aligned} & \underset{\text{Som of } 1 \text{ et } \times 1 = 2 \text{ et } (\text{reduction})}{\text{HgSO}_q} + & \underset{\text{Som of } 1 \text{ et } \times 2 = 2 \text{ et } (\text{existation})}{\text{HgSO}_q} \end{aligned}$$

- Use the multiplier obtained above to balance loss and gain of electrons $H_aSO_4 + ZHI \longrightarrow SO_3 + I_3 + H_4O$
- Balance the rest of equation by Inspection method.
 H_gSO₄ + 2H₁ → SO₈ + I₄ + 2H_gO

NeCl+ MnO₂ + H₂SO₆ \implies Na₆SO₆ + MnSO₆ + H₂O + Cl₆

Identify the elements, which undergo a change in existation number and write their oxidation numbers over the syrabols.

Determine the number of electrons gained and lost and equals them

$$\begin{aligned} & \underbrace{\text{NeCl+ MinD}_2 \times 1 = 2 \text{ c backurion}}_{\text{inne of } 1 \text{ c'} \times 2 = 2 \text{ c (outsition)}} \end{aligned}$$

Use the multiplier obtained above to balance loss and gain of electrons

of Charles of Party

E05

Busines the rest of equation by inspection method

identify the elements, which undergo a change in exidation number and write their oxidation numbers over the symbols.

$$K_0^* C V^* D^* + H_{C_0}^* \longrightarrow KC_0^* + C C^* + H^* D_0 + C^*$$

In this eq. CI is excitised from -1, in HCI to 0 in Cl_q . But it's excitation state is not changed in F/CI I or $CrCl_q$. Therefore, write HCI twose and determine the no-of-electrons gened and lost and equate them

HGI +
$$K_2Cr_2O_7$$
 + HCI \longrightarrow KCI + $CrCl_6$ + H_6Cr + $CrCl_6$ + H_6Cr + H_6Cr

Use the multiplier obtained above to balance loss and gain of electrons $HCI + K_gCr_gO_{\gamma} + 6HCI \longrightarrow KCI + CrCl_s + H_gO + Cl_s$

Belance the rest of equation by inspection method. SHCI + $K_2Cr_2O_3$ + 5HCI \longrightarrow 2KCI + 2Cr O_3 + 7H₂O + 3Cl₃

$$K_{*}C_{7_{2}}O_{7} + 14HC_{1} \longrightarrow 2KC_{1} + 2C_{1}C_{6} + 7H_{6}O + 3C_{6}$$

Chestic Publ

 $Q \in (B)$ Edmant the following equations by in n electron med

Sm⁸⁺ + Fe⁸⁺ -- Sn⁴⁺ + Fe¹⁺

(Surgadin Board, 3019)

Identify the elements, which undergo outdation and reduction and split up the reaction into exidation and reduction half reactions.

Write down the number of electrons gained and lost in each half reaction

$$Fe^{a_{r}}$$
 + $1e^{r}$ \longrightarrow $Fe^{a_{r}}$ (reduction half reaction) _____(1)

Set** — Set** + 2e* (caddedon half reaction) ____(2) Equate the total member of electrons gained and lost by multiplying eq.(1) by 2, and then add them.

$$\begin{array}{ccc} 2Fe^{4} + 2e & \longrightarrow 2Fe^{8} \\ & Sn^{4} & \longrightarrow Sn^{4} + 2e \\ \hline Sn^{8} + 2Fe^{8} & \longrightarrow Sn^{4} + 2Fe^{8} \end{array}$$

$$H^+ + C_1^{-} + C_{12}O_1^{-2} \longrightarrow C_1^{2+} + C_{16}$$

 Identity the elements, which undergo oxidation and reduction and spitt up the reaction into ordination and reduction half reactions.

$$Cr_sO_r \stackrel{s_*}{\longrightarrow} 2Cr^{s_*}$$
 (reduction half reaction)

Balance oxygen by adding H₂O.

$$Cr_2O_7 \xrightarrow{a_1} \longrightarrow 2Cr^{a_2} + 7H_aO$$

 $2Cl^2 \longrightarrow Cl_2$

Belance hydrogen by adding H* lons.

$$14H^{+} + Cr_{z}O_{z}^{-1} \longrightarrow 2Cr^{3+} + 7H_{z}O_{-}$$

Write down the number of electrons gained and lost in each tail read

Equate the total number of electrons gained and lost by multiplying eq.(2) by 3. And then add the two half reactions. 14H1+ + Cr2O2 2 + 6€ -> 2Cr34 + 7H,0

Cu + NO₂ + H⁺ -- Cu²⁺ + NO₂ + H₂O Companies Street, 1990; a identify the elements, which undergo evidation and reduction and upit up the reaction into coddetion and reduction half reactions.

$$NO_a^{-1} \longrightarrow NO_a$$
 (reduction half reaction)
 $Cu \longrightarrow Cu^{2+}$ (oxidation half reaction)

Belance oxygen by adding $H_2\mathbb{O}$.

$$NO_a^{-1} \rightarrow NO_a + H_iO$$

Balance by drogen by adding H* ions.

Write down the number of electrons gained and lost in each half reaction

With down the number of electrons gained and
$$2H^* + NO_8^{-1} + 1e^- \rightarrow NO_1 + H_0^-$$
(2)

Cu -> Cu² + 2e (Z)

Equate the total number of electrons gained and lost by analoghing eq.(1) by 2 And
then add the total number of electrons gained and lost by analoghing eq.(1) by 2 And then add the two half reactions.

When add the two half reactions.

$$4H^{\bullet} + 2NO_3^{-1} + 2e^- \longrightarrow 2NO_2 + 2H_0$$

 $Cu \longrightarrow Cu^{2^+} + 2e^-$

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 $C_{2}O_{3}^{\frac{1}{2}}$ + $H_{4}A_{5}O_{3}$ \longrightarrow C_{2}^{34} + $H_{4}A_{5}O_{4}$

Identify the elements, which undergo oxidation and reduction and split up the Marking into oxidation and reduction half mections.

$$C_2O_3^{-1} \longrightarrow 2C_4^{-1}$$
 (reduction half reaction)

$$H_0AsO_0 \longrightarrow H_0AsO_0$$
 (oxidation half reaction)

Balance cupyen by adding H₂O

$$Cc_{g}O_{7}^{g_{-}} \longrightarrow 2Cr^{g_{+}} + 7H_{g}O$$

$$H_{\bullet}O + H_{\bullet}A_{\bullet}O_{\bullet} \longrightarrow H_{\bullet}A_{\bullet}O_{\bullet}$$

Balance hydrogen by adding H* ions.

White down the number of electrons gained and fost at each half seaction

$$34H^{+} + 6e^{-} + Cr_{2}O_{7}^{2-} \longrightarrow 2Cr^{3-} + 7H_{2}O_{-}$$
 (1)

$$H_{a}O + H_{a}AO_{a} \longrightarrow H_{a}AoO_{a} + 2e^{-} + 2H^{+}_{-}_{-}$$
 (2)

$$261_sO + 311_sAuO_u \longrightarrow 211_sAuO_u + 6e^- + 6f1^o$$

$$Cr_iO_i^{-1} + 8H^* + 3H_iAnO_i \longrightarrow 2Cr^{1*} + 3H_iAnO_i + 4H_iO$$

gate excitation and reduction held reactions.

Charles Service 2013 MnO₃ + C₂O₃ -> Mn³+ CO₃ Identify the elements, which undergo ossidation and reduction and uple up the restriction and uple up the restriction.

$$\begin{array}{cccc} \operatorname{MinO_0}^* & \longrightarrow & \operatorname{Min^{g_*}} & & \{ \text{ reduction hall reaction } \} \\ C_2O_0^{g_*} & \longrightarrow & 2\operatorname{CO}_0 & \{ \text{ oxidation hall reaction } \} \end{array}$$

original Charactery: Post-I

Balance exygen by adding H₂O MnD₄ → Mn² + 4H_aO

$$C_2O_4^{-1} \longrightarrow 200_2$$

G_MHa+**Me ← Gord +*HB

从一届新山村市出口社。1879年

. Write down the murroer of electrons gained and lost in each half reaction

$$C_2O_4^{2-} \longrightarrow 200_2 + 2e^-_2$$

 Equate the total number of electrons gained and lost by multiplying sq.(2) by 5 and eq.(1) by 2. And then add the two half reactions. 10e*+ 16f1*+ 2MnO_e* --- 2Mn²+ SH₂O

$$Fe^{2\alpha} + Cr_2O_7^{2-} \longrightarrow Cr^{2\alpha} + Fe^{4\alpha}$$

 Identify the elements, which undergo oxidates and reduction and split up the mean into ordination and reduction half reactions.

o ordination and reduction half rections.

$$Ce_2O_2^{2-} \longrightarrow 2C_2^{2-}$$
 (reduction half reaction)
 $F_0^{2-} \longrightarrow F_0^{2-}$ (ordination half reaction)

Belance oxygen by adding HyO.

Balance hydrogen by adding H* lone

Write down the number of electrons gained and lost in each half reaction.



Charles Park

(11)

$$14H^{+} + 6e^{-} + Cr_{g}O_{g}^{-1} \longrightarrow 2Cr^{2+} + 7H_{g}O_{----}(1)$$

$$pe^{a+} \longrightarrow Fe^{a+} + 1e^{-}$$
 (2)

Equate the total number of electrons gained and lost by multiplying eq (2) by 6. And then add the two half reactions.

14H* + 6e* + Cr₂O₇** - → 2Cr^{3*} + 7H₈O

$$|O_a^- + AsO_a^+ - \Rightarrow |-+ AsO_a^+ -$$

identify the elements, which undergo oxidation and reduction and split up the reaction into excision and reduction half reactions.

$$AsO_6^{4+} \longrightarrow AsO_6^{4-}$$
 (coddation half reaction)

Belance may gen by adding H_8O .

$$O_3HE + 1$$
 \leftrightarrow O_3HE

$$H_0O + AsO_0^{-3} \longrightarrow AsO_0^{-3}$$

Balance hydrogen by adding H* ions,

$$H_0O + A_0O_0^{-1} \longrightarrow A_0O_0^{-1} + 2H^+$$

Write down the number of electrons gained and lost in each helf reaction

$$H_0O + AsO_0^{3-} \longrightarrow AsO_0^{3-} + 2H^* + 2e^* = (2)$$

then add the two half reactions.

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2H₆O + 3A₅O₆³ · → 3A₆O₆¹ +6H² + 5e.

B. WHITE BEET

Identify the elements, which undergo oxidation and reduction and split up the reaction into oxidation and reduction half reactions.

. Balance oxygen by adding H₂O.

$$7H_{0}O+2Cr^{k_{0}}\longrightarrow Cr_{0}Or^{k_{0}}$$

Billings hydrogen by adding H* lors.

$$6H^* + BiO_3 \longrightarrow Bi^{**} + 3H_2O$$

 $78ig9 + 2C_2^{**} \longrightarrow C_{ip}O_7^{**} + 14H^*$

Wittendown the number of electrons gained and jost in each half reaction

Equate the total number of electrons gained and not by multiplying eq.(1) by 3, and then and the control of the

College Chambrier Part (

612

identify the elements, which undergo ordination and reduction and split up the receipt into oxidation and reduction half reactions.

(reduction half-reaction)

[oxidation half reaction]

Balance caygen by adding HaO.

$$2S_aO_a^{a_a} \longrightarrow S_aO_a^{a_a}$$

Balance hydrogen by adding H* ions.

$$2S_{\alpha}O_{\alpha}^{-1} \longrightarrow S_{\alpha}O_{\alpha}^{-1}$$

Write down the number of electrons gained and lost in each half reaction

$$2e^{-} + OCI^{-} + 2H^{+} \rightarrow CI^{-} + H_{0}O$$
 (1)

$$25_{s}O_{s}^{a-} \longrightarrow S_{s}O_{s}^{a-} + 2e^{-}$$
 _____(2)

Add the two half reactions.

$$25_{2}O_{3}^{3-} + OC_{1}^{2} + 2H^{*} \longrightarrow S_{4}O_{3}^{3-} + Cl^{-} + H_{4}O$$

$$MnO_4 + C_2O_4^2 \longrightarrow MnO_2 + CO_2$$

Identify the elements, which undergo oxidation and reduction and split up the residence of the control of the c Into oxidation and reduction half reactions.

, reduction half reaction)

$$C_tO_t^{2-} \longrightarrow 2CO_t$$

(oxidation half reaction)

Add two OH-tons for one each oxygen atom on appropriate side.

Charlette: Post-

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$$MnO_4^- \longrightarrow MnO_1 + 4OH$$

Balance hydrogen by adding H₀O lons.

$$C_0O_0^{-2} \longrightarrow 2CO_0$$

. Write down the number of electrons gained and jost in each half reaction

$$3e^{-}+2H_{e}O+MnO_{e} \longrightarrow MnO_{e}+40H^{*}$$
 __(1)

$$C_a O_a^{-3-} \longrightarrow 200_a + 2s^+$$
 ___(2)

Equate the total number of electrons gained and lost by multiplying eq.(1) by 2 and eq (2) by 3. And then add the two half reactions.

$$6e^- + 4H_2O + 2MnO_6^- \longrightarrow 2MnO_6 + 8OH(-3C_2O_6^3^- \longrightarrow 5CO_2 + 6e^-$$

$$3C_3O_4^{2-} + 2MnO_4^{-} + 4H_8O \longrightarrow 2MnO_9^{-} + 6CO_8^{-} + 6CH^{-}$$

MnQ₂+ CNU * (D.G. Khen Board, \$009) MnO_s + CN

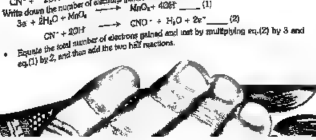
identify the elements, which undergo caldation and reduction and split up the reaction into oxidation and reduction has reactions.

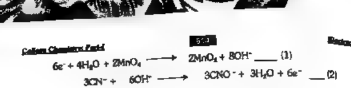
CN° + 20H CN° + H₂O (2)

Write down the number of electrons gained and test in each hell reaction

MnO_x + 4081 (1)

3e + 2H₂O + MnO₄





3CN^+ H₂O + 2MnO₄" ---> 2MnO₂+ 2OH" + 3CNO

 $H_aAsO_a + Cr_aO_a^{a_*} \longrightarrow Cr^{a_*} + H_aAsO_a$

identify the elements, which undergo conduction and reduction and split up the reading into exidation and reduction half reactions.

 $Cr_2O_7^{2-} \longrightarrow 2Cr^{2+}$

(reduction half reaction)

 $H_aAsO_a \rightarrow H_aAsO_a$

, coddation half reaction)

Balance oxygen by adding H₂O.

Cr₂O₇ ²· → 2G²⁺+ 7H₂O

H_BAsO₄ + H_BO --> H_BAsO₄

Balance hydrogen by adding H* lons.

 $14H^{+} + Ct_{2}O_{7}^{2} \xrightarrow{} 2Ct^{3} + 7H_{2}O$

 $H_aAsO_a + H_aO \longrightarrow H_aAsO_a + 2H^*$

Write down the number of electrons gained and look in each half reaction

14H* + 6e* +Ct₂O₇** --> 2Gr** + 7H₂O _____(1)

 \rightarrow $H_iA_6Q_1 + 2H^+ + 2a^-$ (2) H_aAsO₂ + H₂O

 Equate the total number of electrons gained and lost by multiplying eq.(2) by 3. Am then add the two half reactions.

14H+ + 6e++ CrgO, 2+ ---> 2Cr2++ 7H2O

 $3H_aAsO_a + 3H_aO$ \longrightarrow $3H_aAsO_4 + 6H^4 + 6e^2$ (2)

Cr₂O₇ 1 +8H++ 3H₂A₂O₃ → 3H₂A₂O₄ + 2Cr² + 4H₂O

Charlet v. Post-l



Multiple Choice Questions from PAST PAPERS

1 Oxidation No. of oxygen in OF₂ is: (Labore Board, 2016)

2. The highest reduction potential in electrochemical is of F₂ and its value is (Resultable 2012) (c) +2.8V

(b) 2.87V 3. The process during which one metal is deposited on the surface of an other metal by (a) -3.87V using electrical current is called (Revulpted Board, 1909, 2017)

(a) Electrolysis

(b) Electroplating

Delectrolyfic refining (d) Electrolytic purification 4. Stronger the oxidizing agent, greater is the (Recolpted Secret, 2010, 2012; Gapren in Secret, 2014, 2014)

(a) exidation potential

(b) reduction potential

(d) E.M.F of cell

5. The electrolyte used in fuel cell is: (9.5) Man Board, \$220; Labora Board, \$200; D.G. Alter Board, \$200; Gulfranzola Board, \$200

(a) acqueous NaCl (b) mother NaCl The oxidation number of chronium in E₂C₂C₃ is: (Exception Board, 2009) (Interest 2009, 2010)

16, 12 7: Oxidation state of chromina in NaCiQa is a shore Beams, sais)

(c) 3 (d) 7

8. Neison's cell and Down's cell are examples of (Outremote Board, 2002; Palacided Secret, 2003)

(a) electrochemics cell (b) galvenic cell (c) electrolytic cell (d) none of these

9. In reaction 3Fe + 3Cls - 2FeCls, regiments are all (d) none of these
(a) Fe is reduced (b) Fe is outdised or Cls is outdised (d) none of these (d) none of the longitude of the recharged are called: (Behandly Sand, 2009 Gelement of the Recharge of the Re

(b) secondary cells (c) tentiary cells (d) primary cells

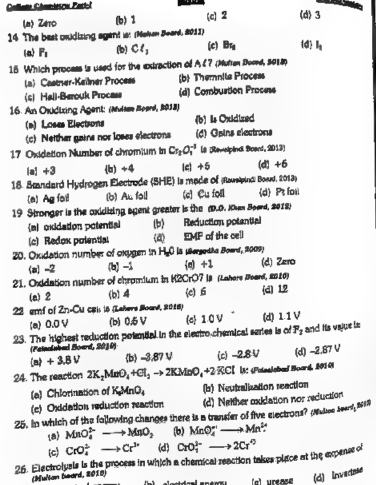
(a) circusta cens (b) character of capper in responsible found, 2012) (d) -2

(c) +2 (a) +1. 12. Electrode potential of SHE entitoerity taken in voits in (Outree) * (d) 0.50

(a) 0.00

(a) 0.00

13. Oxidetion number of an element in free state in: grades flours, 2011.



(b) slectrical energy

(a) Chemical energy

(e) urease

Chemina Paris g in conversion of Br_2 to BrO_3 the oxidation state of browning changes to go LOS (b) 0 to -3 (c) This 5 (e) 0 to -2 S. According to classical concept, oxidation involves suggests some some (a) Addition of oxygen (b) semoval of hydrogen (e) increase in oxidation sinte (d) all of above 29. Bulphur has highest exidation state in (Submodyor Found, 2010) Encourage Sures, 2010) (b) SO₂ (c) H₂S [d] H₁8O₁ (p) 8O₃ 30. Oxidation state of carbon in glucose is (0.0. Khim Beens 8810) (b) one (c) two (a) 2010 31, Which one is not an electrolyte (Lakers Board, 2017) (a) Aqueous NaC1 (b) Aqueous CuSO, (c) Cumetal (d) H.SO. 32. Oxidation state of Mn in MnO 3 is platon food, 2011) (a) +4 (b) +633, Cathode in NICAD cell is (Pateriphed Board, \$10.1) (d) 2a (a) AgeO (b) NiO₂ (c) Cd Act of Report, 2012) 34. Percentage of H₂SO₄ used in lead accumulator is ## (d) 50% (a) 40% (b) 25% (c) 30% 35. Loss of electrons is called (Labora Bound, 2007) (a) Oxidation (b) reduction (c) hydration (d) dehydration 36, Pupi cells convert chemical energy into (Labora Bourt, 2007) (a) Heat energy (b) Light energy (c) Electrical energy (d) Mechanical energy 37. The coddation number of Mn in KMnO₃ is Grandwood Second, 2027) (a, 3 (b) 5 (c) 7 (d, 9 38. If salt bridge is not used between two half cells, then the voltage order (a) decreases rapidly
(b) decreases slowly
(c) does not change 39. In Na₂O₂, the oxidation state of oxygen is Geografia Goard, 2011 (a) -2
(b) +2
(c) +6
(a) zero
(b) +6 (a) zero

610 ton Charleton Peri-Answers to Multiple Choice Questions from Past Papers. Ans Q# Q# Ans Q# Am Ans Q# Ana tal Б. 3 (b) 4 [c] 2 (c) (c) 9 **(b)** 10 (c) 6 (d)7 (b) 6 (c) 14 16 13 (a),c, 12 (a) (b) 11 19 20 (b) 18 <u>(b</u>, (a) 17 (d) 16 <u>(a)</u> 24 25 (c) (b) (d) 23 22 (c) 21 (c) 29 30 (a) 28 (d) (a) 27 (c) 26 (b) 35 34 <u>(b)</u> (0) <u>,a)</u> 33 (b) 32 91 (c)

Detailed Explanation of Past Papers MCQs & Answers to all Past Papers SHORT QUESTONS In COLLEGE CHEMISTRY OBJECTIVE BOOK-I

38

(c)

 (d_I)

39

40

(a)

SHORT & LONG QUESTIONS FROM PAST PAPERS

OXIDATION NUMBER, BALANCING OF EQUATION BY OXIDATION MUMBER AND TOR ELECTRON METHODS

Short Quastions

36

(c)

(1) Define electrochemistry. (February)

37

- (2) Define oxidation with an example (Guarannia Bount, 2011) Sorg
- (3) Define reduction. Give an example (Nature Board, 2007)
- (4) Differentiate between exidation and reduction, Calors Board, 2002 (5) What are redox reactions? Give example. (Guyranuda Board, 2012) rd, 2008; Paleda
- [6] Define the oxidation number with an example. (5=gotha Board, 2016)
- [7] Glue Rules for axidation Number of Hydrogen and Oxygen. (Suspolite Sc
- (B) Prove that the oxidation number of some elements vary in different compounds (Sepadas Board, 2007)
- (9) What is the oxidation state of CI in KCIO and KCIO₃ (D.O. Khen Beard, 2009)
- Colculate the oxidation number of N in HNO, and NO. & material Board, 2011
- Calculate addation numbers of manganese in KMnO₄ and sulphur in SO₄.
- Colculate the exidation number of phosphorus in: (Revelpted Board, 2018) (a) H.PO, (b) No.PO.
- Lene Questions

 [1] Define oxidation number Describe the rules used for the colculation of activity. number of an element in molecules and ions. (Recording beard, 2007; Schauelper Beard, 2007).
- (2) Give steps for balancing the Radox equation by Oxidation number method. **

Name and Address of the Lot

CONDUCTION, ELECTROLISMS, ELECTROLISMS PROCESS OF GROWINGAL PROCESS bert Chrestions

3.75

- (i) Define (a) electrolytic conduction (b) Electrolytic call Define (c) Define electrolytic giving an example, required the second state of the second secon

- Differentiate between electrolysis and electrolysis conduction (Residual and series)

 (4) What will be the products and reactions of the electrolysis of makin (or hand) NoCP d' Board, 2009) (Gajramada Garri, 2014)
- [5] Write a brief nate on electrolysis of aqueous solution of NaCl, and Write docum cathodic and anodic reactions involved in electrolysis of squeous solution of sodium chloride. (D.G. Kim Bourd, 2000) On Give Coshodic and Anodic reactions of electrolysis of concentrated aqueous solution of Sodium Chieride photos to
- (6) Glue two applications (or uses) of electrolysis processes of industrial importance, glabasosper general, 2009; D.G. Khar Bourd, 2011. Makes Bourd, 4616, 1019

 17) What is the industrial importance of electrolysis? plumpata theret, 2018

- Loss Guestians (1) Describe the electrolysis of molton NoCl and a concentrated existing of NoCl statement, 2009: Galyonnada Scard, 2009: Sugardia Band, 2011; Later Scard, 2017 (Septimbrie) Band, 2017; Sugardia Band, 2018; Later Scard, 2017; Sugardia Band, 2018.
- Food, 2009: Gujronanda Board, 2009; Sargecha Brand, 2015; Labor Josef, 1915 (2) Situe the Industrial Importance of electrolysis process in detail? 1944; Materialand Board, 2015)

SALVANIC CELL, ELECTRODE POTENTIAL AND ITS MEASUREMENT, STANDARD, INDROGEN ELECTRODE

- (1) What is standard electrode potential? One econode principal Band, 2009, 2019, 2019. Des Notes Bend, 2009: Recompised Seard, 2011: Volum Band, 2019. Descripted Band, 2019: Laws Band, 2016: Gajimmada Band, 2014) [2] What Is S. J. F. P. Give its value, (Bahamdan Band, 1989) [3] One S. J. F. P. Give its value, (Bahamdan Band, 1989)
- II) OR What do you mean by standard hydrogen electrode for SHE? What is in unitOR What stands for SHE? What stands for SHE? Define it. (Separal Bond, 2007) February Bond, 2009. Labora Bond, 2010)

 State (or Differentiate) oxidation and reduction mactions. (Selection Bond, 2012)

 State (or Differentiate) oxidation and reduction mactions. (Selection Bond, 2012)

- What is the function (or purpose) of sak bridge in growth calls, planning from 1912 2012, 2013; East, 2019, 2013, 2014. Recorded Sound. 2019 Sept. 2019, 2013, 2014. Recorded Sound. 2019 Sept. 2019, 2013, 2013. Recorded Sound. 2018 Sept. 2019,
- How can the say that a voltate out is reversible call dis-
- Salvanic cell? (Sargodia Souri, 2009) Write anacht and eathers Reactions of Generic cell. (1999)

Chambios: Park!

Charles de la constitución de la

Long Grantons

(1) Describe the construction and working of standard hydrogen electrode, (Sugadha Board, 2015); Revelyted Board, 2014)

2005: Outcomed Board, 2013: Laters Board, 2015; Revelyted Board, 2014)

(2) How can you measure electrode potential of an element with the help of standard (2) How can you measure electrode potential of an element with the help of standard (2) How can you measure electrode potential of an element with the help of standard hydrogen electrodes. How our you measure encurses Board, 2012) OR What Is standard hydrogen electrode (SHE)? (Latera Board, 2012) OR What Is standard hydrogen electrode hydrogen electrode joints. I have the electrode potential of zinc? (Lakora Board, 2014) On (SHE)? How it is used to measure the electrode potential of zinc? (Lakora Board, 2014) On (SPE). How it is used to measurement of an unknown electrode potential with the help What is S.H.E.? Glove the measurement of an unknown electrode potential with the help

425

of it. (Between Beard, 2010) (3) What is electrode potential. How electrode potential of Zn is measured? (Malson Box

2011)

(4) Define electrode potential. Describe the construction of voltaic cell and reaction occurring In the cell. (Labors stoard, 2009) OR Describe a galognic cell explaining the function of electrodes. (Remaintable Beard, 2009)

ELECTROCHEMICAL SERIES AND ITS APPLICATIONS

(1) Define electrochemical series. Give its one application. (Behaviolpus Board, 2009; Labort Board, 2007, 2015; Sargodia Board, 2019) NOTE: For one application see page (2) How can use calculate voltage of a cell with the help of electrochemical series. Give

example. (D.G. Rhan Board, 2012)

(3) What is emf of a cell? (Nutren Bo ard, 2007)

Write two applications of electro-chemical series. (Guirnovala Board, 2013)

(5) Describe the relative chemical reactivity of metals (Bahas four Board, 2011) OR Host electrochemical series inelp us in determining the chemical reactivity of metals? (Guja

(6) Differentiate between artificing and reducing agent. (Supremuta Board, 2008)

Long Oversions

(1) What is electrochemical series? Also explain two/three applications of electrochemical series. (D.G. Kleen board, 2007: Gapranosis Spord, 2613; Sargodius Sourd, 2013)

PRIMARY AND SECONDARY CELLS

(1) Define primary cells and secondary cells (Sergodia Board, 2009)

(2) What are primary cells? Give example, Ofiction Board, 2007)

(3) Differentiate between a primary and a secondary cell. (D.C. Hass Board, 2004: Multon Ba

(4) What is the difference between a cell and a battery? (Mutua Board, 2009; Bull-2010; D.G. Khan Bourd, 2010)

LEAD ACCUMULATOR, ALKALINE BATTERY SILVER OXIDE BATTERY

Short Questions

(1) Give electrode reactions in lead storage battery. (Fesseleted Board, 2009, 2015)

(2) OR Describe redex reactions which occur during discharging of lead accumulator batter).

(Supposts Board, 2013: Malten Boa

Continue Paris

671

grant is Lead Accumulator? Discuss in detail (Makes Board, 2016) on Exploin "Lead Accumulator" with changing and discharging, so a Rose Board, 2016 on Exploin "Lead What is Levin recharging and discharging, and share Board, 2016 on Explain "Len Accumulator" with charging and discharging, and share Board, 2017 Familiand Board, 2018 when the silver Oxide battery. OR Write a rate on ellipse of the board, 2018 (4) Describe the silver Oxide battery OR Write a rate on silver oxide battery. Re-

out Overtions (i) Write down the reactions in allowine battery. Oblice Bows, 2009 OR What are electrode

NORE CADMIUM BATTERY, FUEL CELLS

What is NICAD? (Gujna nto Bound, 2010) OR Writz down reactions taking place at the What is treatment assume the discharging of Nickle-Cadmium Cell, itakes 80rd, 8021)

electrones us are some new trial and where it is used? (D.G. Minn Board, 2007; Ramaphal Board, 2009)

[3] Gipe chemical reactions taking place at anode and azinode in a fuel cell (labor hand)

How power is generaled by using the fuel cell? (Revelated Board, 2007)

15) Where are fuel cell used? How much fuel bond energy is concerned into electricity in them? (Bahawalpur Board, 2008)

(b) Give some advantages of fuel cell. (Labore Board, 2011,

Long Queeltogue

(1) Describe Nickel Cadmium Cell (rechargeable) (Labora Board, 2014)

[2] Write a note on the construction and working of fuel cells. (Asset Kasts

[3] Explain finel cell with its construction, electrolytic reaction and diagram (separal)

Write comprehensive makes on alkaline battery and nickel-coamium battery. (Salas Board, 1912)

Marks: 68 SUBJECTIVE Time: 2:10 Hours Note: Cut of Questions 2,9 and 4, Write any TWENTY TWO(22) short answers. While writing and (22x 2) =44 quarties numbers carefully. Section - [

Of. Answer may Eight parts from the followings.

(a) Cu

(f) Write down the electrode reaction of eliver oxide bettern.
(ii) Zn can displace Cu from CuSO₄, but Cu can not displace Zn from 2n5O₄. Why?

govern reducing agents have high value Second measured experies have high well What are the functions of said bridge? What is modified As ? in of standard reduction po What is encountered of an electrolytic cell, in which electroles also be example of an electrolytic cell, in which electroles also be exact to Flati and Bercult process? if the one occurred to an account of the state of the control of t How does electrochemical terior tell ut the distinction between the control of th How toes excupation in a chargeable benery, justify to principals between conduction and reduction parameter of the princip cells? Give one example laster my Eight parts from the following in the first printerly cease Give one example

of Assert any Eight partie from the followings.

Stranger as anode when connected to Du theringle but as rathods with 2n election.

How does oxidation reduction reaction to be place during declaration conduction?

What is injustical importance of electrolytics?

What reactiffs products of electrolytic activities of PlaC. Give reactions?

What care of Colvania could be reversed?

What are callede and anode?

What is SHE? self What is SHE?

(c) When Zu is compled with SHE, then which reaction doors at 2n electrode?

(c) When the reaction Ca2* + Zu Cu + Zu⁴ is sportaneous or not? Give reacon?

(d) What is insert by, smit of a Galversic call?

(d) Abalt metals (L., Ne, is and Rb) are highly reactive and coinage metals (Cu. Ag and An are seen.) | Placi means the same the followings. |
| Why? |
| Why? |
Why	Place	Place	Place
Why	Place	Place	Place
Why	Place	Place	Place
Why	Place	Place	Place
Why	Place	Place	Place
Place	Place	Place	Place
n) Give Rules for conduction Number of Hydrogen and Onygen? Ni-Wate anodic and cathodic reactions of Gavanic cell?			
M) What is NICAD? M. White in NECAD?

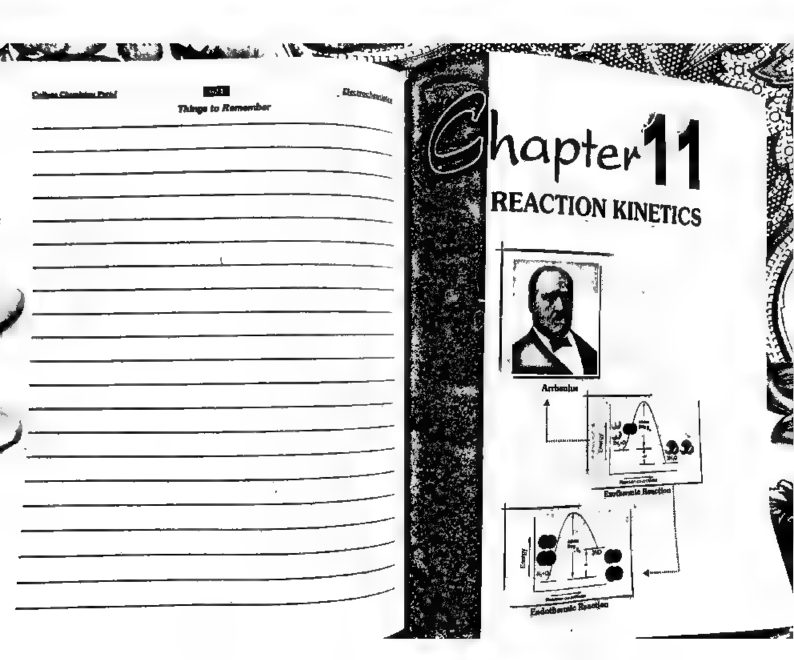
M. Name three chemicals which are used as firsts in fuel edits?

All Wall is meant by the term caude tion number or state of an elem

(a) White is the outdeddon state of Cr in K_BCr_BC₇? (5 is) What is electrochemical series? Give its three applications. (94) (a) Describe the process to determine the electrode polential amagonic. (b) Describe the process to determine the electrode polential area more reactive? Explain with reference to electrodemics. (c) Determine a fuel cell generate electricity? (c) Determine the extra born state of Cr in KCO₀ and O₀O₀ (c) with down the oxidation state of Cr in KCO₀ and O₀O₀ (c) with down the rules for assigning exclusion states to elements in a compound. (b) Describe the construction and working of a Celeptic cell. (02)

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at Chemistra Pend



CONTENTS -

Chapter-11 -

Reaction Kinetics

INTRODUCTION

RATE OF REACTION

Instantaneous and average rate Rate constant or velocity constant. Order of reasons

Half-life period or half life time

Rate determining step

DETERMINATION OF THE RATE OF

CHEMICAL REACTION

Раумскі теблобі Constructed mathenda

ENERGY OF ACTIVATION

FINDING THE ORDER OF REACTION

Half life method

Method of large excess

FACTORS AFFECTING RATE OF

REACTIONS

Nature of reactant

Effect of concentration

Surface area

Lagist

Effect of temperature

Catalyet CATALYSIS

Characteristics of satalyst

Activation of catalyst or promoters

Enzyme catalysis

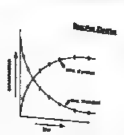
Characteristic of enzyme catalysis

Objective and short suggest questions (exercise)

Past Papers MCOs and Short Onesteens

Test your skills

Office Contains Part I



TO CAS

STRODUCTION

The branch of chemistry, which deals with the study of reaction rates and the futu-shich affect those rates, is longers as chemical kinetics or reaction invites. Reaction kinetics helps to determine the mechanism of vactions.

Sei

THE SOL CHEMICAL REACTIONS

On the basis of velocity of reactions, there are three types of reactions.

the Reactions

Some reactions occur very slowly and take even years to complete. e.g., Consider the reaction

This riskellon occurs very alowly and might take, perhaps, decades to complete.

Strallar are the nusting of iron, chemical weathering of stone work of buildings by addic Pass in the atmosphere, tennentation of sugas etc.

in t Rom there

Some reactions take aimost no time to complete. Such reactions are last reactions.

e.g. Consider the reaction

This reaction is completed as soon as AgNO New and NaClog louches each other.

deal ante Reactions

Some reactions occur at moderate rates, neither two slow nor too feet.

^{e.g.} Hydrolysus of enter

CH°COOH + C°H°OH ---- OH°COOC°H° + H°O

Tali Stop Reactions

Por multistan reactions, one of the step is slow. This is called rate determining step.

Other steps do not affect the rate of seaction.

ALCOHOLDS ROBERTS

to the other billions are the

District to characters suspicion, proclama and composited into participate.

the reaction tenouries the epocational of reacting decreases with time and the responsibility of product increases with time. Therefore, make of seaction can be defined as

The value of shareness to come Applied of Persons

on The spin of increase in any contention of pro-

When reaction starts, concentration of mactants decreases with time, while concentration of product increases. At start, rate of meadlion is generally very fast. Thus, the slope of care is greened for both reactants and products as shown as the fig.

After some time, the slope becomes Jess steep showing that motion rate <u>decreases</u> with time. It shows that rate of a seaction is changing at every moment,

Actually, pair of mection is decreasing with time. It never pins uniform during different time period. It decreases continuously until the seaction steps. gen i de la symmette describité de la la La activides and producte, pathilise



Usually, concentrations are expressed in mol dm⁻⁸ and time in sec. Therefore, units of cate are

Rule =
$$\frac{Ac}{At} = \frac{\text{mol}/\text{dm}^2}{\text{sec}} = \text{mol} \, \text{dm}^{-3} \, \text{sec}^{-4}$$

For asseous reactions, certial pressures of asses are used instead of molar concentrations.

INSTANTANI OUS AND AVERAGE RATE

SVERIAL HATE

The rate of a spection between epocific time interpole is called the non-3s rate of

Let amount of product formed at t_i is ϵ_i and at t_j is ϵ_k , then

Average rate =
$$\frac{c_3-c_1}{t_2-t_1}=\frac{\Delta c}{\Delta t}$$

MANIANEOUS EATH

Pet X, to the success of inogens harmony as not gas I, the I approved as

The Article (chillerential) of x w.r.t. Y) is called min of change of once west force to y enternis instantaneous reaction une

decard all one very school changes in concentration and time impuribility For a general seaction

The rate of meactions can be expressed in terms of tale of disapplinance of A or use of appearance of B i.e.

Rate =
$$-\frac{d[A]}{dt} = +\frac{d[B]}{dt}$$

Whene 'd [A]' and 'd [B] are very small change is concentrations of A and B moectively.

Negative sign undicates that conc. of A decreases with time, while Positive sign undicates that conc. of B increases with time.

39 Moeship herricen Instantaneous out Accrese Resea

- At start, instantaneous rate is greater than average rate
- At the end of the interval, the instantaneous rate-becomes year than average mee.
- When the time interval approaches zero average and indestructure nine become equal.
- Average and instantaneous rates become equal for only one instant during the Interval.

Par Constant or Velocity Constant

- Rate constant is the rate of reaction when the conscious of reactions are using
 - It is denoted by 'k
 - Under give conditions, k remains constant
 - It changes with temperature

beforetten.

According to the law of Mass Action

have of a chemical reaction is directly proportional to the product of active masses of and the product of active masses of edants involved in the chemical reaction.

Thus for a simple reaction aA + bB -----→ products

Rate = $k(A)^a (B)^b$

Where k = specific rate constant

When [A,=[B],=1 mol/dm² then

Rate = k

Thus, Rate constant is the rate of reaction when the concentrations of reactions are units

ORDER OF REACHON

The number of reacting molecules whose concentration changes during a reaction is celled order of reaction

Consider a general reaction

aA + bB ----- products

Rate equation for this reaction will be

Rate of reaction = k[A]*(B)*

a is the order of reaction w.r.t A

b is the order of reaction w.r t 'B

The sum a + b is called overall order of reaction

Thus, order of reaction can be defined as

The sum of the powers to which concentrations are related in the rate for expression is called order of reaction

Order of a reaction is an experimental quantity it cannot be determined by reaction

In chemical kinetics, reactions are classified as zero ender, first order, second order and third order

Zero Onler Reaction

When the reaction is independent of the concentration of reactions, it is called simorder reaction.

Examples

Decomposition of [4] (on gold surface)

2HI on gold region Hg + Ig

The experimental rate equation for this reaction is

Rate = $k(H!)^{0} = k$

This reaction is independent of the concentration of HI, hence it is a zero order reaction. Photochemical reactions are usually zero order.

Charleto: Part-I

when each of powers to which concentrations are retaid in the rate less cope pol to 1.

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Samples.

1 Decomposition of nitrogen pentoxide.

 $N_2O_4 \longrightarrow N_2O_4 + i/_2O_2 \longrightarrow$

The experimental rate equation for this reaction is

Rate = $k[N_sO_s]$

Hence # Is first order w.r.t. the concentration of N.O.

2. Hudrolysis of territory butyl bromude (Preudo Pine Order territor) The reactions in which vate depend upon only one molecule diffough more than on plecules are impolived is called a pseudo first order reaction.

tertiary butyl bromide

ĊH,

lartiery butyl alcohol

The experimental rate equation for this reaction is

Rate = $k ((CH_a)_5CBr)$

Hence, it is first order w.r.t. [CH₂]₁CBr. The rate of reaction is independent of the transmitted of wester, because, it is present in very large success. Thus, it is a passeto that order reactions.

Second Order Renetions

When sum of powers to which concentrations are raised in the rate less expression is क्षात के हैं.

temple

Catation of nitric oxide with ozone

$$NO_{\omega} + O_{0 \omega} \longrightarrow NO_{2 \omega} + O_{1 \omega}$$

The experimental rate equation for this reaction is

This reaction useflest excises when NO and first order with O. The sum of the individual takes is equal to two. Hence it is a second order reaction

Hard Order Rometions

When sum of powers to which concentrations are roteed in the rate lass expression to equal to 8

Example

Reaction of FeCl, with KI

This reaction involves eight reactant molecules. However, the experimental rate equation for this reaction is

Rate =
$$k [FeCl_j] [KI]^k$$

This reaction is first order w.r.t. FeCi, and second order ω r.t. KL

The sum of the individual orders is equal to three. Hence it is a third order reaction. It shows that it is a multistep reaction and one step is the rate determining step. The possible steps are

$$\begin{array}{lll} \text{FeC}_{\text{s} \text{ tot}} + 2 \text{KC}_{\text{loc}} & \xrightarrow{\text{dem}} & \text{FeI}_{\text{l} \text{ tot}} + 2 \text{KCI}_{\text{loc}} + \text{CC}_{\text{loc}} \\ 2 \text{KC}_{\text{loc}} + 2 \text{CC}_{\text{loc}} & \xrightarrow{\text{loc}} & 2 \text{KCI}_{\text{loc}} + 1_{\text{l} \text{ tot}} \end{array}$$

NOTE:

The order of a reaction is usually positive integer or a zero, but it can also be in fraction or can have a negative value

Example: Formation of Carbon Tetrachlaride Form Chloroform.

The experimental rate equation for this reaction is

Rate =
$$k \left[CHCl_{g} \right] \left[C_{lg} \right]^{1/6}$$

The sum of exponents will be $1+\frac{1}{2}=1.5$

Hence the order of this reaction is 1.5.

The experimental relationship between a reaction rate and the concentration of the reactants is known as the rate loss or the rate equation for that reaction.

Consider a general reaction

Rate eq. for this reaction will be

Characters: Part

Rate = k[Af |B]b

'a and 'b' are distermined expertmentally. Rate law expression is, therefore, an sperimental expression

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PRI THE PERIOD OF HALF LIFE UME

the time required to convert 50% of the reactants late products is called half by of the

It is denoted by the or tag

people 1: Decimoposition of N₂O₄

 $2N_2O_{1 \otimes i} \longrightarrow 2N_2O_{1 \otimes i} + O_{2 \otimes i}$

Halfille period for decomposition of NgO₈ is 24 minutes at 45°C

However, that if reaction is started with $0.1 \, \text{mol/dm}^2$ of $N_e O_0$ then after 24 number, 0.06note/dm⁵ will be left behind

Similarly, after 48 minutes (two half life periods), 0.025 mol/dm² (25% will be left behind and after 72 minutes (three half life periods), 0.0125 movium (12.5%) will be with behind Decomposition of N₂O₄ is a first order reaction

Above temperiment clearly shows that half life time of a first order reaction is indimendent of the initial concentration of reactants.

Exercise: Disintegration of 350

It has a half life period of 7.1 x 10° years or 710 million years. It is also a first order

It megans that if 1 kg of $^{235}_{92}\mathrm{U}$ is present, then after 710 million years, 0.5 kg is converted this daughter elements, and in the next 710 years, 0.25 kg of the 22 will be consumed.

defutition of the section of Reaction

Generally, How the period is inserestly proportional to initial concentration of reactants raised to function one less than the order of reaction.

$$(t_{1/2})_n \propto \frac{1}{a^{n-1}}$$

n = Order of reaction a = Initial conc of reactantsMpalé. $\{t_{i/2}\}_n = Half life period for nth order reaction.$

College Chambirly Part |

Reaction States

• For first order reaction, half life period is independent of the initial concentration of

Let
$$\{t_{1/2}\}_{i} \propto \frac{1}{a^2}$$
 , since $\{t_{1/2}\}_{i} = \frac{0.693}{k}$

a For a second order reaction, half-life period is inversely proportional to the high concentration of reactants,

Let
$$\{t_{1/2}\}_2 \propto \frac{1}{a^2}$$
, since $\{t_{1/2}\}_2 = \frac{1}{kn}$

For a third order reaction, half life period is inversely proportional to the suitage of Initial concentration of reactants.

Let
$$(t_1/p)_3 \propto \frac{1}{a^3}$$
, since $(t_1,p)_3 = \frac{1.5}{k_B^2}$

Hence half-life period can be used to determine the order of reaction.

Example I

date the half-life period of the following respection when the initial cane. Is 0.05 M

$$2HI_{\rm tot} \implies H_{\rm tot} + I_{\rm tot}$$

$$k = 0.079 \text{ dm}^2 \text{ mol}^{-1} \text{ s}^{-1} \text{ at } 508^4\text{C} \text{ and } \text{Sate} = k[H])^2$$

k = 0.079 dm^b maj -1 s -1

concentration of HI = a = 0.05 M.

Since the reaction is second order, therefore

Per a second order resiston

$$(t_1/2)_2 = \frac{1}{k \times a^{2-1}} = \frac{1}{k \times a}$$

$$(l_{1} \ z)_{2} = \frac{1}{0.079 \times 0.050} = 258 \frac{1}{600}$$

College Chemistry Part-I

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Esercise Q11.

Exercise Less.

The rate determining step of a reaction is found out from mechanism of that reaction Explain it with few examples. Take second example on page 643, 2NO + 2H₂ → 2H₃O + N₃

WATE DESIGNATION STEP

The reaction, which determines the operational of reaction, is called rate

For multistep reaction, rate determining step is the slowest step. Employation.

Reactions may occur in single step or several steps. The reactions, which occur in more than one sime, are called multistep reactions.

For a single step reaction, the only step is aways the rain determining step.

For multisten reactions. One of the step is slower. This is called rate determining step. Other steps do not affect the rate of reaction.

The total number of reactant molecules that takes part in rate, determining step episons In the experimental rate equations of reaction.

Higgies, experimental rate solutions can give information about machinism of quantions. Emmole 2:

The experimental rate equation for this reaction is

Rate =
$$k [NQ_b]^2$$

This rate equation shows that rate is independent of concentration of CO. Thus, it shows that if Is a multistep reaction Morsover, one step is slow which is the rate-determining Map and involves two molecules of $N\!O_{\rm f}$

Thus, following mechanism can be written

$$NO_2 + NO_3 \xrightarrow{\text{this}} NO_1 + NO$$

So, first step is alow and it is the rate determining step. NQ_{ij} is not produced in the reaction, it is consumed as soon so it is formatic. Such a Specie is called reaction intermediate.

Repution Internadiate

a

The specie which produce only for abort period of time during a reaction to collect

Reaction intermediate is unstable we considered to both renderate and products. In wome bases, under certain conditions, it may be stable and can be looked.

with regular intercals

2111(g) H₁(t) + I₂(g)

0.100

0.0716

0.0558

0.0457

0.0387

0.0336

0.0296

0.0265

0.0

Exercise Q10

Exercise Q10

A curve a obtained when graph is plotted between time on x-axis and concentration on y A curve a commen union graph on the slopes of various points gives us the instantaneous rates of reaction. Explain with suitable examples.

DESTRUCTION OF THE WATER TO THE MICAL WEACHERS

While measuring rate of reaction, progress of reaction is noted gither by physic methods or chemical methods. In these methods, conc. of one of the reactant or product a determined at different time intervals.

The different concentrations obtained are then piothed against the time. The slope of curve at any time I' on conclume graph will give the Rate of Reaction at that time.

The slope at any point on the graph (e.g., at point P) can be obtained by drawing tangent to the curve at that point. A night angle triangle ABC is completed with znomi as invoctenuse, such that

Rateof reaction= $\frac{AC}{BC} = \frac{\Delta C}{\Delta t}$

Exemple

Consider the decomposition HI to H₂ and I₃ at 508 °C

The conc. of Hil determined at various times is given in the table

This table clearly shows that for first 50 sec, the decrease In conc. of HI is 0.0284 mol/dm3, while between 300 to 350 sec, the decrease is only 0.0031 mol/dm³

Hence, rate is fester at the start, but slower at the end

It has also shown by the slope of graph. Graph shows that slope is greater at the beginning than at the end. And greater the slope higher is the rate

The rate of this reaction at any time can be determined by drawing tangent to the graph he required time. Shows of the determined the graph of the determined time. at the required time. Slope of the tangent will give the rate at that time.

e.g. at point P,

slope = rate = $\frac{0.027}{140}$ = 2.5×10⁻⁴ moles dm⁻³s⁻³ 110



50

ION

150

250

The night angled biangle can be of any size, some tale of reaction will be obtained

If a graph is plotted with concentration of H_{ϵ} or I_{ϵ} a tising r is re will be obtained However the rate at any time will be the same as with graph or

EXPERIMENTAL TECHNIQUES FOR MEASURING RATE OF REACTION Physical Methods

In physical methods, progress of reaction is noted, by observing the changes in physical properties of the reaction mixture.

Most community used physical methods are

ii. Spectrometry

or Chembers: Part-1

(ii) Conductometry (iii) Dilatometric method

(iv) Retractrometry (v) Polarimetry

Spectrometry

This technique is used if reactant or product absorb LIVIVIS or IR light

The amount of radiation absorbed is directly proportional to the progress of reaction

Conductometry

This method is best for <u>logic reactions</u>. Each ion has its own <u>conductivity</u> when ions are formed or consumed during a reaction, then conductivity chances. Hence progress of reaction can be noted by measuring the conductivity changes

The conductivity change is directly proportional to the rate of change in concentrate the ions.

Dilatometric method

This method is used for reactions in which vokums is changed.

The volume change is directly proportional to the progress of reaction

Refractionetes

This method is used for reactions in solution, in which refractive indices of substances The change in refractive index is directly proportional to the progress of reaction change

Optical Rotation Method (Polgrimetra)

This method is used when one of the substances is anticelly active. ...e. if can retain the Diane polarized fight. The angle of rotation is determined by polariquete

The change in angle of rotation is directly proportions, to the progress of reaction.

After measurement, graph is plotted and rase of reaction is determined from the graph.



According Manhouse to these methods, samples are withdrawn from reactions vessel at different time interest In these methods, stropped are was assumed by in order to determine the concentration of the of the reaccasts or products at different faces.

3.53

Consider the hydrolysis of ethyl acetate

OHCOOCH+HO === OHCOOH+CHOH

During this maction samples are taken out of reaction. vessel at different times and placed into about four times of its volume as an cold water to stop the reaction. The diamon and chilling stop the reaction. The concentration of CityCOOM produced in the reaction is noted every time by timeting against standard alkali solution using phenolphthalten as indicator

The different concentrations of acetic acid are plotted against the time. A rising curve is obtained, initially rate of Xseaction in high but it decreases with time. The mile is determined from the concentration-time graph. When the <u>curve becomes hortents</u> the pate becomes zero.

If curve is plotted with <u>concentrations of searchast,</u> then a <u>falling curve</u> is obtained

it is a <u>psyculo first order</u> reaction. Since water is present in large excess as compare to ^{est} therefore, it does not affect the rate of reaction.

INERGY OF ACTIVATION

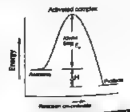
cules must have, in addition to the rage K.E., to form on

it is denoted by E.,

Units: It is expressed in joules

In a reaction system, constantly collide with one another

At ordinary temperature, activation energy is usually very high for any reaction system. Therefore, most of the molecules



6.59

supply relogand after collisions and do not form activated complex

However if some of the molecules gain enough energy after collisions to form activated complex, they will start the reaction. Such collisions, which form an activated complex are called effective collisions. For effective collisions, molecules must have proper orientation.

if all the collisions are effective then the reaction will be completed soon, ricure, a most of the reactions are slow showing that all collisions are not effective.

The number of effective collisions increases with increase in importating and thus rate of eaction is also <u>Increased</u> and vice versa.

Colliding molecules first form a high energy transition state or activated complete, which then leads to product. It can be represented as

Activitied complex is an unstable, short-fived intermediate, it at once changes into product.

When colliding molecules <u>come</u> close to each other, they show down, collide and fit spent. If the collision is <u>effective</u> then the <u>fitting molecules</u> will be <u>different</u> if not, then same molecules will bounce back. When the molecules above down just before collision, then K.F. decreases and P.E. increases.

Relationship Between all And En-

The heat of reaction is equal to the difference of potential memo, between reactants and

Exotherade resultan

For exothermic reactions, greater of martent is picture, then the spears of product. The difference of energy is released as heat. The Secresse in P.E. of the residents appear as increase in K.E. of the



It has been shown in the fig.

Carlotherm/o repetitori

For endothernic reactions, energy of machine

is lower than the energy of product. The difference of energy is absorbed as hase. Such reactions, a continuous supply of energy a needed.



It has been shown in the fig.

It is clear that, both for exotherwic and endothermic reactions, there is energy harrier, which must be overcome in order to start

For all reactions, activation events is different for forward and backward reactions. For exothermic reaction, E. Is less for forward reaction than backward seaction. And for endothermic reaction. E a more for foreset then between reaction. E give resource about guidensess of reaction.



PARSO THE ORDER OF REACTION

Many methods are used to detectains the order of reaction

- His and Tree method.
- Geographical method
- Differential positions
- Hall life snethod
- Hethod of large recent

Juli-life method

y proportional to initial concentr than the order of reaction.

$$(a_1/g)_a \propto \frac{1}{a^{a-1}}$$

Where n = Order of reaction

a. = Initial conc. of reactants

 $\{|t_{\rm LR}|\}_a \approx Mail life period for <math display="inline">n^{th}$ order reaction.

Let a reaction is carried out with two different initial concentration of reactants a Let two half-life times are to and to respectively.

Then

$$t_i \stackrel{\circ}{\alpha} \frac{1}{z_i^{n-1}} = (1)$$
 and $t_i \propto \frac{1}{z_i^{n-1}} = (2)$

Divide eq. (11by (2)

$$\frac{h}{t_2} = \frac{1}{2} \frac{1}{a_1^{n-1}} = \left(\frac{a_2}{a_1}\right)^{n-1}$$

Taking log on both sides

$$\log\left(\frac{t_1}{t_2}\right) = \log\left(\frac{de}{dt}\right)^{n-1}$$

or
$$\log \left(\frac{t_1}{t_2} \right) = (n-1) \log \left(\frac{a_2}{a_1} \right)$$

$$\frac{\log\left(\frac{h}{t_2}\right)}{\log\left(\frac{a_2}{a_1}\right)} = \{n-1\}$$

$$n = 1 + \frac{\log\left(\frac{t_1}{t_2}\right)}{\log\left(\frac{a_2}{a_1}\right)}$$

Thus knowing $t_{\rm g}$ and $t_{\rm g}$ at $t_{\rm g}$ and $t_{\rm g}$, order of seaction in item be determined.

in the thermal decomposition of N₂D at 760°C, the time required to decompose holf of the reactions uses 255 sec, at the initial P of 290 sun Hg used 212 sec at the billial P of 250 mm Hg, Pind the order of this reaction.

Folutions

For 1st experiment

initial conc. \approx a, ≈ 290 mm

Half life time = $t_1 \approx 255$ sec.

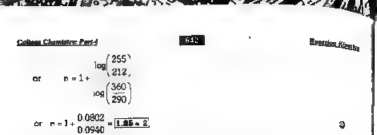
For 2⁻¹-experiment

Initial cone. \Rightarrow $a_{ij} = 360 \, \mathrm{mm}$

Half life time $= t_0 = 212$ sec

Order of praction is given by

$$m = 1 + \frac{\log\left(\frac{1}{t_2}\right)}{\log\left(\frac{m_2}{t_1}\right)}$$



Hence it is a second order reaction.

Method of Large Excess

in this method, one of the reactants is taken in small amount while others are taken in large amounts.

Since cond. of reactants present in large amount is almost constant. Therefore, rate-doss not depend on their concentrations

Hence rate only depends upon the cond of reactant taken in small amount. It is because small changes in its amount greatly affect the rate. Thus order of reaction wirl this readant is determined

Similarly, this process is repeated with all the reactants by taking them in small amounts Thus order of reaction w.r.t. each reactant is determined.

Hence overall order of reaction is the sum of the orders wirit, each reactant.

Exercise O12:

Discuss the jactors influencing the rates of chemical reaction.

FACTORS AFFECTING RATE OF REACTIONS

Following factors affect the reaction rates

- 1. Nature of reactant
- 2. Concentration of reactants
- 3. Surface Area
- 4 Light
- 5. Temperature
- 6. Catalyst.

(I) NATURE OF THE REACTANT

Reaction rates are greatly affected by the nature of Reactant The reactivity of a substance is controlled by its electronic arrangement.

College Charlette: Part-

Examples

· ·

Elements of troup I—A reacts more rapidly with H₂O than elements of troum II A.

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- Neutralization and double decomposition reactions are very fast.
- Jonic reactions are very fast e.g. reaction between HCI and NeOH is completed in Just 10⁻⁶ sec, at room temperature,
- · Redox reactions involve transfer of electrons, therefore, these are slower than sonic

2: EFFECT OF CONCLNIRATION

Bate of reaction is affected by changing concentration in accordance with law of Mass Agiton. This law states

The rate at which a substance reacts, is directly proportional to its notice mass, and the rate of reaction is directly proportional to the product of active masses of reacting

Thus by increasing cone, of any reactant, rate of reaction is increased and by decreasing cond, rate of maction is decreased.

The <u>frequency</u> of <u>collisions</u> between molecules depends upon their <u>concentrations</u>. The <u>proper crowded</u> the molecules, the <u>more chance of collision</u>. Hence, <u>rate</u> of maction increases.

Examples

- Combustion in normal air (with 21 % $\mathbb{Q}_{g'}$ becomes faster in pure O_0
- Lime stone reacts with different rate with different concentration of HCI
- Concentration of pages can be increased by increasing their pressure and hence rate of reaction increases.

Exercise Q18.

Explain the following facts about the reaction

The jointning jets about the recomm. $2NQ + 2H_2 \rightarrow 2H_2Q + N_1$ $2NQ + 2H_2 \rightarrow 2H_2Q + N_1$ The changing concentrations of readons change the roles of this reaction. Individual orders with respect to NQ and H_2 can be measured. Individual orders with respect to NQ and H_2 can be measured. industrial angles with respect to the order to the concentration of one of the The overall orders can be evaluated by keeping the concentration of one of the

꼢 subscances constant

Consider the reaction

Excerimental data for this relation is given below

College Chambers, Part-L

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Reaction Reputing

(1 totale (1 1.2) Effect of change in concentrations of reactants un the rate of reaction in

(NO) in (profite day 1)	(1(_j) is (males din ⁻²) = ¹ 22	i Initial rate (afin mina)
Ö 006	0.001	0,025
0.006	0.002	0.050
0.006	0.003	0.075
0.001	B.009	D.0063
< 30%	0.009	0.025
70%	0.009	. 056

Experiments show that by keeping concentration of NO constant, and doubling the we entry ion of $|\underline{f}_{b}|$ \underline{double} the read ion <u>rate</u> and <u>tripling</u> the concentration of H_{b} <u>triples</u> the

Thus rate of reaction depends on the first power of concentration of H₂

e Rate a [Ha] +1,

Similarly, keeping purcentration of He constant and doubling concentration of NO. raises the rate to 4 times, and inpline concentration of NO raises the rate to 9 times.

Thus, rate of reaction depends on square of concentration of NO

Combining 1) and (2)

Rate a [Ha] [NO]

Rate = $k |H_1| |NO_1^{\dagger}$

It's the rate law for the given reaction. It shows that it is a third order reaction. Thus, rate equation for this reaction is not according to balanced equation

To satisfy rate eq. following mechanism can be proposed

$$2NO + H_R \xrightarrow{\text{then}} N_1 + H_1O_2 \text{(nate determining step.)}$$
 $\tau_0 U_3 + H_4 \xrightarrow{\text{finit}} 2H_1O$

3) SURFACE AREA

By receasing a risce area, contact between specified molecules increases, tience paged reaction increases.

Example.

- A tall reacts stoods with NaOH but powdered Al reacts rapidly with cold NaOH * 2NaAi-OH_{/4} + 3H₆ 2A - 2NaOH + bHO
- Cal O3 in big pieces react stoods with H2SO4 but it reacts rapidly in powdered form

Orders Chemistry Part-I

H HGHI Light consists of photons of definite enemies. When light a given to a reaction, its enemy ncreases the rate of reaction Examples

- Reaction of CH₄ and Cl₄ requires light
- Reaction of H_g and Cl_g is pagilistic in darkness, slow in dark light but exclusive in sunlight.
- Photosynthesis requires light.
- Reactions of silver halide in chotography requires light

5) Effect of Temperature

According to collision theory, rate of reaction is proportional to the number of collisions and the malecules. Thus, if increasing of collisions is among the molecules. increased, rate of reaction is also increased. However all the collisions do not result in reaction. Only effective collisions can

give activation energy to the molecules.

To start reaction, molecules must rain activation energy E.

At low temperature mostly molecules posses average energy and only small fraction of molecules posses necessary E, for reaction as shown in the fig by shaded area. Most of the molecules possess average energy

With increase in temperature, the traction of molecules with high energy negatives. The curre of higher temperature T, has lightened. Thus, there is which temperature T, has lightened.

distribution of velocities. At higher importance T₀, number of molecule having high energy discreases while number of molecules with low shared decreases. Thus, effective collisions will be a lowered decreases. be increased and the rate of reaction will also be increased.

When the temperature is reised by 10 K, the fraction of moreouse with energy more than E. Dughly doubles. Thus, the rate of reaction is also doubled

Q15 How does Archechus equation help us to celculate the mergy of antivertion of croction?

Arrhenius Femation (Effect of T)

Increase in 7 increases the sets of rection and also set Generally, rate of reaction doubles for every 10°C rise in temperature. Effect of ismperature on reaction rate it best explained by Arrheticia equation

 $k = Ae^{-6680}$ (1)

Where k= rate constant

A = Frequency factor or Arrhenius constant

e = Base of natural logarithm. Its value is 2.718281

E. = Energy of Activation

R = General gas constant

T = Absolute temperature

'A' depends upon the collision frequency of the reactants.

Taking natura; log of eq (1)

$$tnk = tnA - \frac{E_n}{RT}tne$$

Since $\delta n = 1$ and $\delta n = 2.303 \log$, therefore

$$2.303 \log k \approx 2.308 \log A + \frac{E_a}{RC}$$

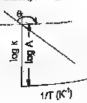
Divide throughout by 2.303

$$\log k_i = \log A - \frac{E_s}{2.303 \text{ M}} \qquad (2)$$

This equation shows that

- Higher the activation energy, lower will be rate constant
- Higher the temperature, higher will be the rate constant.

Intercept = log A



Calculation of Activation Forces

Eq (2) is an equation of straight line.

Le
$$y = -mx + c$$

Where 'm' = slope of line

c' = intercept of straight line

A graph between $\log k$ and $\frac{1}{T}$, willigive a straight line with

Slope =
$$\frac{\hat{E}_b}{2.2020}$$
 and intercept = log A

Thus Slope =
$$\frac{-E_a}{2.303R}$$

or $E_a = -stope \times 2.303 \times R$

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Slope is determined by measuring tapacet of and θ . Angle θ is obtained by drawing a line perallet to x - axis and measure angle 6 Take tan 9 which a the

Different reactions will have different straight lines and different shows

Units of Slope:

Slope=
$$\frac{E_0}{2,303R} = \frac{J^2 \text{mole}^{-1}}{2,303J \text{mole}^{-1} \text{K}^{-1}} = K$$

A plot of Arrhenius eq. for the thermal decomposition of $N_0 D_0$ to choose in the foll $N_0 D_0$. The alope to found to be -5800 K. Calculots the energy of activation of this reason.

Amhenius eq. is



From Anthenius eq.

This shows that the thermal decomposition of NrO4 requires 103:39 μ_0 moi $^+$ energy in

addition to the average energy or maximit to cross the energy barrier



6) Caralesi

A substance that changes the reto of reaction without being consumed in the reaction

A catalyst increases the rate of reaction by decreasing the activation energy of $t_{\rm lg}$ reaction.

Example

 Reaction between H₂ and O₂ is very slow at room temperature. However, it speeds up in the presence of Pt catalyst

CATALYSIS

Cotalest

ce that changes the rate of reaction but remains chomically unch the end of reaction is called catalyst,

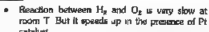
Function Of Catalyst

A catalust lowers the activation energy of the reaction by providing a different way for reaction as shown in the fig

By lowering E, more molecules can easily cross the energy barrier and hence reaction rate increases.



of London Section (1997)



- mom T. But it speeds up in the presence of Pt catalyst
- KCIO₃ decomposes much more rapidly in the presence of a small amount of MnO₃
- HCI is exidized to Ci₂ in the presence of $CuCl_2$ 4HCI + O_2 $\frac{cucl_2}{2H_2O}$ + $2Cl_2$

CATALYSIS

The process, which takes place in the presence of a entalyst, is called outsiyets.

A catalyst gives a new reaction path with low activation energy

Thus more molecules can cross the new energy barrier and reaction rate increases

Types of Catalysts

- ,1) Homogeneous Catalysis
- (2) Heterogeneous Camiyais

College Chemistre: Pari-I

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(1) Happingeneous Catalesis

In this process, the catalyst and the reactants are in the a

The reacting system is homogeneous throughout and the catalysi is distrib through the system.

Examples:

1. Oxidation of SO₂ to SO₃ in the lead chamber process for the manufacture of sulphuric acid, needs NO as a catalyst.

Both the reactants and the catalyst are gases.

2. Esters are hydrolysed in the presence of H₂SO₄. Both the reactants and the catalyst are in the solution state.

 $\mathsf{CH_3COOC_3H_{0,\mathrm{col}}} + \mathsf{H_2O_{0}} \xrightarrow{\mathrm{chi_3COOH_{\mathrm{loc}}}} \mathsf{CH_3COOH_{\mathrm{loc}}} + \mathsf{C_2H_2OH_{\mathrm{loc}}}$

$10 \text{ ns } pns pn_1 = 0$

In this process, the establet and the reactants are in different phases.

Mostly the catalyst is in the solid phase, while the resciants are in the g_{θ} phase.

Emmoles

1. Oxidation of ammonia to NO is carried out in the presence of platinum gazes during manufacturing of HNO₃.

Hydrogenation of unsaturated argenic compounds are catalyzed by finely divided Ni

3. Oxidation of SO_{ϵ} to SO_{ϵ} in the contact process for the manufacture of sulphwise acid, needs V₂O₆ as a catalyst.

CHARACTERISTICS OF A CATALYST A calculate not changed in mass and chambrai camposition of the end of reaction.

e.g. For the decomposition of KGO_3 , MnO_4 is added in the form of granules. It is convented to fine powder at the end of reaction its uhysical state may be changed. In many cases shining surfaces of the solid catalyst become dulf. 2. Sometimes only a hard of a mind counts is used to affect year larger amount of



Reserve Karney

Exemples

- 1 mg of fine platinum powder can convert 2.5 dm 3 of H $_2$ and 1.25 dm 4 of $_{\rm C}$ 0,
- Dry HCl and NH₃ combine in the presence of trace of mousture to give design white furnes of NH₄CL
- Φ Thousands of dm³ of $H_{\rm g}O_{\rm g}$ can be decomposed in the presence of 1 g σ colloidal platinum.
- 3. A catalyst is more effective in finely divided form, it is because increase in safety area increases the efficiency of a catalyst and rate of reaction increases. e.g. a big piece of Pt have much less calalytic activity than colloidal Pt. In the hydrogenation of vegetable oils finely divided nickel is used.
- 4. A capabest connect affect the equilibrium constant of a reaction. If only decrease the time to reach equilibrium.
- A catalyst cannot start a reaction, which is not thermodynamically feasible.
- 6. The mechanism of a catalysed reaction is different from that of an uncatalysed
- 7 Specificity: A calabat is specific in its action. A particular catalyst works for one reaction. It may not work for any other reaction.
- B. If different catalysts are used for the same renotant then the products may charge. Examples

Formic acid is decomposed by Al₂O₂ to water and CO while Cu decompose if to H_z to CO_z.

HOOCH -Ca + Hz + CO2

- Affect of Temperature: Temperature affects the role of a catalust. Physical state of some catalyst is changed by change in T, Hence their catalyst power will be decreased, e.g. colloidal catalysts like platinum may be coagulated with the rise in temperature.
- 10 Poleomina of a Cotolivet: Catelian may be poleomed by trace amounts of other substances. Thus they become ineffective. Such substances are called poisons. The poisoning of a catalyst may be temporary or permanent. In permanent poisoning, the poison reacts chemically with the catalyst Example: Compounds of sulphur and arsenic act as poisons to many metalic catalysts. e.g. Pt catalysts, e.g. Pt

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Escuphes

- Decomposition of H₂O₂ is cassived by colloidal platform. Pt can be made ineffective
- (ii) In Haber's process, presence of CO as an impairing with hydrogen decreases the
- (iii) In Contact process of manufacturing of H₂SO₄, Pt is used as a catalyst. Traces of arsenic as impurity in the reacting gases makes PI anethoctive. That's why assenic

Activation of Catalyst or Properties

A substance, which promotes the activity of a catalyst, is called a pre-

It is also called "catalust for a catalust".

Examples:

- † Hydrogenation of vegetable oils is carried out by Nt. The catalytic activity of Ni can be increased by using Cu and tellurium,
- ii) In Haber's process for the manufacture of animonie, from is used as a catalyst. If anall amounts of some high-malting orders like aluminum catale, chrorekum exide or rare earth orders are added, the efficiency of iron is increased.

NUGATIVE CATALAST

A substance, which decreases the role of reaction, is called a negative or labilities.

Ехотиче

Tetraethyl lead is edded to petrol, because it saves the petrol from pre-tyridion.

In some of the reactions, a product formed may set us a catalyst. This phononemon is called auto-catalysis.

Camples

- i Provincitais of which accesse generales scalle acid, which access as a catalyst for further 2. When popper is allowed to cent with next ecid, the reaction

Colon Charles Fat-I

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Specific Paris

beginning, it gains speed slowly and finally becomes very fast.

Despiriture, a genus special property of nitrous acid during the reaction, which special to be

 The reaction of oxalic acid with acidified KMnO₆ is slow at the beginning but MrSQ. produced in the reaction makes it faster 2KMnO₄ + 3H₂SO₄ + 5(COOH)₂ K₂SO₄ + 2MnSO₄ + 10CO₂ + 8H₂O

Exercise Q18.

What are enzymes? Give examples in which they act as catalyst. Mention the charactering of enzyme autalysis?

LNZYML CATALYSIS

rms are the complex protein molecules and catalyse the organic reactions in the dis. They are called blochemical entelpots.

Many enzymen have been identified and obtained in the pure crystalline state. The fin enayme was prepared in the laboratory in 1969.

I. Uses undergoes hydrolysis into NHs and CO_2 in the presence of ensyme asset

2. Glucose is convented into ethanol by symase ensyme present in the yeast.

3. Concentrated sugar solution undergoes hydrolysis into glucose and inucles $^{\rm tr}$ invertage exceptive, present in the yeast

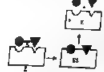
$$C_{12}H_{22}O_{11} + H_2O \xrightarrow{\text{Long.}} C_2H_{22}O_4 + C_4H_{22}O_4$$

Enzymes have active century or cavities on their surfaces. The molecules of a substitute form country of a substitute form. fit into cavities, form complex, reacts and the product is get out of the cavity immediately.

Callery Chamistry, Parts

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Michaulis and Menter proposed following mechanism for enzyme case



Characteristics of <u>Fazenas Catalesis</u>

The role of enzymes as catalyst is like morganic historogeneous catalysts. They are highly efficient and specific

- 1. Enzymes are the most efficient catallets known and they lower the energy of activation of a reaction
- 2. Erroving catalysis is highly specific
 - e.g. <u>urease</u> catalyses the hydrolysis of uses only and it corner hydrolysis any other arnide even methyl urea.
- 3 Enzyme catalysic reactions have maximum rates at an optimum temperature.
- The phi of the medium also controls the rates of the ensyme comit, sed reactions; Each enzyme has an optimum pH.
- 5. The activity of enzyme is greatly enha-

ion is independent of the

Since A is in exame, therefore, sale will depend of conc of B only Harres, it will be a first order rach

Per Ares (d)
According to the collision theory, rate of dependent upon the number of collision reaction molecules With increase in temps number of collisions among reacted mole increases. Hence, rate of nacrice doublet.

a Rate = k*[B]

	A -> Products
	If this resident is a zero creder erection, then its rain equation is given as
	Risks = K(A) ⁿ =ic Prence units of the thereony see that to rote of maction. (a. mot drawns ⁻¹)
	The state of machine Let mod dis-
đs.	Fill in the blanks with suitable words.
	(ii) The rate of endothermic reaction with the increase in temperature (iii) All radioactive disintegration reactions are of order (iii) For a fast reaction the rate constant is relatively and half-life
	(iv) - The second order reaction becomes if one of the reactants is in far excess
	(v) Arthenius equation can be used to find out of a reaction.
	Answers
	All Increases All files and All generalized for first order tel energy of arthration
CMI.T.	adicate TRUE or FALSE = the case may be.
Sacial	(i) The half life of a first order reaction increases with temperature.
	(ii) The reactions having zero activation energies are instantaneous
	term a site of the second of t
	 (iv) There is no difference between the area of the state of the confer of reaction is strictly determined by the statehomenty of the balanced equation
_	
	Answers. fit Polse
	A language with physical
Q4	What is observing kinetics? How do you compare chambon kinetian with chambon equilibrium and thermodynamics?
Chair	troad forwattor. It is the study or rates of reachions and the factors, which affects these rates the
-	or raises of reactions and the passess
•	It is the study or rates or temperature, catalyst etc. concentration, temperature, catalyst etc. It gives information about reschenism of reactions. It gives information about reschenism are studied in chemical kinetics.
	It oftens information about measure and studied in chamilton renerce.
	concentration, compared to reactions. It gives information about mechanism of reactions are studied in chemical Minetics. Both reversible and irreversible reactions are studied in chemical Minetics.

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Chambion Peri-l

ret Auss to t

equation is given as Rate = IdA)²=0. It means that the risk of m

ATIS (18)
According to the time of mass action, mail of a repotion,
is climicity proportional to the product of concentrations
of maccaning Air the product of concentrations
concentration of maccaning the concentrations.



Conclus Manager

Chamical equilibrium

- · It is the state of a reversible reaction at which the rate of forward reaction becomes equal to the rate of reverse reaction.
- It is only concerned with reversible reactions.
- Le-chartiver's principle is used to study the effects temperature pressure are concentration on equilibrium state.

Thermodynamics

- It is the study of energy changes.
- It tells about the feasibility of a chemical reaction

The rate of chamical reaction with respect to products is written with positive sign, but with respect to reactants is written with negative sign. Explain II with reference to the following hypothetical receitors.

During a chemical reaction, reactants are converted into products and their concentrations change. Thus, rate of a reaction can be expressed with respect to any reactant or product

However, as the reaction proceeds the concentration of reactants decreases with the and the concentration of products increases with time. Thus, rate of chemical reaction with respect to products is written with positive sign but with respect to reactions, it is written with negative sign

* Hence, rate of above hypothetical reaction can be expressed as

Rate
$$=$$
 $\frac{1}{a}\frac{d(A)}{dt} = \frac{1}{b}\frac{d(B)}{dt} = \frac{1}{a}\frac{d(C)}{dt} = \frac{1}{d}\frac{d(D)}{dt}$

What are the instantaneous rates and average rates? Is it true that instanta rate of a reaction at the beginning of reaction is greater than the sperage rate side of a reaction at the beginning of reaction is greater than the sperage rate section is greater than the sperage rate near the completion of reaction?

The rate of reaction between specific time intervals is called the average rate of reaction tile While

The rate at any one instant during the interval is called instantaneous rate

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Differentiate between

(1) Rate and Rate constant of a reaction

	Mortical factorial form	W	Restrict entertaint of a transferen
1	It is the change in concentration of reactants or products divided by the time taken for the change.		It is the rate of a reaction when the
2	It changes with time.	2	If does not changes with time.
3	It depends upon the concentration of reactants	3	It is independent of the concentration of reactants.
4	It is a variable quantity		[(is a constant quantity.
5	lis units are mol dim ³ 5 ¹		its unit depends upon the order of reaction.
6	Example. Consider a general reaction A + B + Its rate is given as Rate = k[C][O], where 'k' is the rate con-		

(B) Homogenous catalysis and Heterogenous catalysis

	(Sargodho Board, 2011. Ravalpindi Board, 201	3. Gultmings has a
	Aconogramos calcievis .	Wedenships cutaces is
ı	in this catalysis, reactants and catalyst ere in the same phase	in this catalysis, reactants and catalyst are in the different phase
2	In this, the system remains homogenous during the	In this, the system remains heterogeneous during the reaction
3	reaction. Mostly liquids and gases are used as homogeneous catalysts	Mostly soild are used as heterogeneous catalysis.
ı	Executivis: 250 _{top} 250 _{top} (Example: 250 _{max} + O _{tex} state 250 _{max} + O _{tex} state 250 _{max}
4	NO is homogeneous cabalus	11

Justify the following statements

The rate of a chamical reaction is an oper changing parameter under the st conditions.

On What happens to the rate of a chamical receiven with the possage of time?

[Multan Board, 2011 | Labore Board, 2012] (Surgodha Board, 20) 1. Mulian Board, 2012,

According to the law of mass action, rate of a chemical reaction is directly proportional to the concentration of reactants.

When the reaction starts, the concentration of reactants is high, therefore rate of reaction is fast. As the concentration of reactants is decreased, the rate of a reaction is the decreased. At the end of reaction, the reaction becomes very slow. Hence, rate of a chemical reaction is an ever changing parameter

(8) The reaction rate decreases every moment but the rate constant 'k' of the reaction is a constant quantity, under the given conditions. Justify it.

According to the law of mass action, rate of a chemical reaction is directly proportions to the concentration of reactants.

Consider a general reaction

A + BC + D

its rese is given as

Rate = k[A)[B], where 'k' is the rate constant,

As the reaction proceeds, concentration of reactants is decreased, therefore, rate is also decreased. However, rate constant 'k' is not changed. It is proportionality constant.

(iii) 80% of a hypothetical first order reaction completes in one hour. The remaining 50% needs more than one hour to convert itself into products.

The time required for half of the reactants (i.e. 50%) to convert into products is called half tife time of the reaction. For a first order reaction, half life time is a constant quantity under given conditions.

For the above hypothetical first order reaction, helf life time is one hour. Therefore, if we start with 100% reactants then the reaction will proceed as

Turn (hour)	0	1	2	3	4_
Abramit fol Han jaid	100	50	25	12.5	6.75

Hence, after every one hour, amount of reactant becomes half. It shows that 50% of the reactants are consumed in one hour. In next hour, 50% of the reactants are consumed in one hour. In next hour, 50% of the reactants are consumed in one hour. remaining amount is consumed. Thus in a total of two hours, 75% of the total amount consumed

Hence, 50% reactants are consumed in one hour but the remaining 50% will take ment hours to be consumed.

College Chamistry: Part-I

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(bd) The radioactive decay is always a first order rescriber, (Recalphal Board, 2007 D.C. Rhan Board, 2007 Lahore Board, 2007 Feliababad Soard, 2009 Sargadha Board, 2009; Chimasala Board, 2011. 2012, 2014: Sargaciho Board, 2007, 2012: Multon Board, 2012)

The rate of radioactive decay depends on the amount of radioactive substance. Since only one substance (reactant) is involved in this process, therefore, it is always a first order seaction. Moreover, the half-life time for the radioactive decay of a perticular substance is also constant. Hence, it is a first order process.

(a) The traits of rate constant of second order reaction is defined in but the unit of rate of reaction to mol don's'

Rate of reaction is the change in concentration with change in time. Therefore, its units are given as

Rote =
$$\frac{\Delta n}{\Delta t} = \frac{\text{mol}_{x} \, \text{den}^4}{\text{and}} = \text{mol} \, \text{den}^{-4} \, \text{sec}^{-4}$$

However, for a second order reaction the rate of reaction is directly proportional to the concentration of two reactants e.g., A and B. I.e.

Rate=k[A][B], where 'k' is the rate constant.

Since the units of rate-mol dra "sec" and concentration of A and B are expressed in mol dm 4, therefore, units of a will be

$$k = \frac{Rote}{(A|B)} = \frac{rholdm^2 e^{rt}}{rand dm^2 \times mol dm^2} = dm^4 mol^2 t^{-2}$$

(st) The sum of the co-efficients of a balanced chemical equation is not necessity. Superstant to give the order of resolins.
(Superstal Bord, 2009: Superstal Board, 2009: Motor Board, 2012)

A balanced chemical equation only gives the stouchiometry of the reaction. It does not deal with the mechanism of the reaction, However, order of reaction is determined experimentally, and it depends upon the mechanism of reaction. Actually, it depends upon the slow step of the reaction. Therefore, the co-efficients in belanced chunked equation has the valeties and the reaction. no relation with order of reaction, e.g., consider the reaction $2H_k + 2NO \rightarrow 2H_kO + N_k$

Rate of this reaction is

Hence, its order is 1+2-3 (third order months).

Hence, its order is 1+2-3 (third order months) equation=2-2-4.

However, the sum of co-efficients in balanced chemical equation=2-2-4. Sometimes, the sum of co-efficient may be equal to the order of reaction.

(vit) The order of a reaction is obtained from the rate expression of a reaction and the rate expression to obtained from the experiments.

Order of reaction depends upon the machinism of section. Actually, it depends upon Order of reaction depends upon the machinism of section.

College Chambers: Part 1

05/0

the slow step of the reaction, which controls the overall rate of reaction.

→

Consider a general reaction

A+B

C + D

Rate expression for this reaction will be

Rate =
$$k[A|^{4}|B|^{6}$$
.

'a' and 'b' indicates the number of molecules involved in the slow step of the reaction These are determined experimentally Rate expression is, therefore, an experimental expression. Hence, order of reaction (i.e a+b) is also an experimental quantity

Explain that half life time mathod for measurement of the order of a reasion on help us to measure the order of even those reactions which have a fredition

Using half life method, order of reaction can be determined by the equation

$$n = 1 + \frac{\log\left(\frac{t_1}{t_2}\right)}{\log\left(\frac{g_2}{g_1}\right)}$$

where n=order of reaction

 t_{i} is the half life time when the concentration of reactants is a_{i} , and t_{i} is the half life time when the concentration of reactants is a_2 .

Thus knowing to and to at an and as respectively, order of reaction in can be determined The answer can be in whole number or in fraction. Hence, half life time method can be used to determine both simple and fractional order

Q10. A curve is obtained when groph is plotted between time on scale and concentration on y-axis. The measurement of the slopes of various points along at the instruments and a state of the instruments of the slopes of various points along at the instantaneous rotes of reaction. Explain with autable examples.

Solved on Page 636

Q11. The rote determining step of a reaction to found out from mechanism of the reaction. Explain it with face examples.

Solved on Page 635

Q12 Discuss the factors influencing the rates of chemical reaction.

Solved on Page 642

Coffee Chapters: Part 4

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Q15. Explain the following facts about the reaction. 2NO + 8H₂ ->

 $2H_{1}O + N_{2}$

The changing concentrations of reactions change the rates of this re individual orders with respect to NO and H, can be measured.

The operall orders can be seeducted by keeping the concentre

Salved on Page 643

Q14 The collision frequency and the proper orientation of molecules are una conditions for determining the proper rate of reaction. Justify the statement.

Solved on Page 638

Q15. How does Arrhenius equation help us to calculate the energy of action reaction?

Solved on Page 645

Q16. Define the following terms and give examples.

(t) Homogeneous catalysis Solved on Page 649

(II) Heterogeneous catalysis Solved on Page 649

Salved on Page 651 (III) Activation of a catalyst

(to) Auto catalysis Solved on Page 651 Solved on Page 650 (v) Catalytic poteoning

(of) Engrue catalysis

Q17 Briefly describe the following with exemples

(I) Change of physical state of a cotalest at the end of reaction.

The physical state of a catalyst may be changed at the end of reaction.

Example 1: For the decorposition of KCIO, MrO, is added in the form of granules. It is converted to fine powder at the end of reaction.

Example 2: In many cases shaing surfaces of the softi consist become duf.

(10) A very small amount of a casalyst way proce sufficient to carry out a reaction.

Sometimes only a trace of a meta catalyst is used to affect very large amount of

reactams. Example 1. 1 mg of fine platinum powder can convex 2.5 $m dm^2$ of $H_{\rm f}$ and 1.25 $m dm^4$ of

Exemple 2: Thousands of deal of H₂C₆ can be decomposed to the pre-

coloidal piatinum.

(III) A finely divided outsiyst may proce more effective.

A catalyst to more effective in finely divided form. It is because increase in surface and increases the efficiency of a catalyst and rate of reaction increases,

Exemple 1: e.g. a big piece of Pt have much less catalytic activity than colloidal Pt. Example #1 In the hydrogenation of vegetable alls finely divided nickel is used.

(in) Squillierium constant of a researchie reaction is not changed in the presence of a

A catalyst cannot affect the equilibrium constant of a reaction, it only decrease the time to reach equilibrium.

Example: $CH_sCOOH + C_sH_sOH \Leftrightarrow CH_sCOOC_sH_s + H_sO$ In this reaction, few drops of H_sSO_s are used as catalyst. So, the equilibrium is established within hours. However, if catalyst is not used then it may take days to reach the equilibrium in both cases, the equilibrium constant will be same at some temperature.

(Surgodko Boerd, 2007: Lahore Board, 2009)

(a) A catalyst is specific in its outlon. Of A particular catalyst is suitable for a particular reaction. Ja

(Maltin Beard, 2006; Pelsalabed Board, 2011)

A catalyst is specific in its action. A particular catalyst works for one reaction. It may not work for any other reaction.

Example:
Unesse enzyme catalyses the hydrolysis of usen only and it cannot hydrolyse any old enside even methyl urea.

What are enspecied Give examples in which they act as entalpst. Mention the characteristics of enspire entalpsis?

Solved on Page 652

BIPORCANT HOMGLAS

$$n = 1 + 1 \frac{\log\left(\frac{h_1}{h_2}\right)}{\log\left(\frac{h_2}{h_1}\right)}$$

is = Ac-East 46 log k = log A 2.303RY

$$\frac{E_{s}}{\text{slope}} = \frac{E_{s}}{2.309 \text{R}} \quad \text{er} \quad E_{s} \approx -2.309 \times \text{R} \times \text{slope}$$

College Chambers: Part-I

0.62

NUMERICAL PROBLEMS (Caercise)

Q19. In the reaction of NO and H₁, it was discreted that agriculturally relative of pures at 340.5 mm pressure uses half changed in 162 seconds. In another experiment with an initial pressure of 285 mm of H₂, the reaction was half completed in 140 seconds. Calculate the order of receion.

For 1st experiment

Initial concentration $= a_1 = 340.5 \text{ mm}$

Half tife time $= t_1 = 102$ sec.

For 2nd experiment

unitial concentration $\Rightarrow s_2 = 258 \text{ mm}$

Hinti life time $= t_i = 140$ sec.

Order of reaction is given by

or
$$n = 1 + \frac{\log\left(\frac{h}{r_{p}}\right)}{\log\left(\frac{82}{e_{1}}\right)}$$

or $n = 1 + \frac{\log\left(\frac{102}{140}\right)}{\log\left(\frac{288}{340.5}\right)}$
or $n = 1 + \frac{0.1376}{-0.0727}$
 $n = 1 + \frac{0.1375}{0.0727} = \boxed{1.99 = 3}$

Hence, it is a third cader reaction

Q.20. A study of chambial together of a reaction

--- produces

A+8 -

on the following date of \$5°C. Colo



Beaution Street,

LAps.	[6]	181	Hapter \$
₫.	I	0.15	4.2 × 10-4
被	2	0.15	8.4×10 ⁻⁶
496	1	0,2	5.6× 10-4

Colution:

Comparison of Exp. $\{1\}$ and $\{2\}$ shows that Keeping [8] constant and doubling A_i doubles the reaction rate

Thus Rate a [A]I

Comperison of Exp. (1) and (3) shows that keeping [A] constant and changing [B] from 0.15 to 0.2, rate changes from 4.2 \times 10 $^{\circ}$ to 5.6 \times 10 $^{\circ}$

Rate a [B]^L

Hence overall order of rate is given by

Rate a [A]1[B]1

Rate = k [A] [B]

Thus overall order of reaction is $1 \div 1 = 2$

Some reaction toking place around room temperature have activation energies around 50 kJ mot-1

(1) What is the value of the factor $q = \frac{E_{\star}}{RT}$ at 25°C.

 $E_a = 50 \text{ kJ mol}^{-1} = 50000 \text{ J mol}^{-1}$

T = 25°C + 273 = 298 K

R = 8.314 5 mol *1 K*1

$$e^{-\frac{8000}{1314\cdot300}} = e^{-20.181} = 1.72 \times 10^{-9}$$

(III) Calculate this factor at 35°C and 45°C and note the increase in this factor for each 10°C rise in temperature.

Solution: At 3FC

 $E_a = 50 \text{ KJ mol}^{-1} = 50000 \text{ J mol}^{-1}$ T = 35°C + 273 = 308 K

College Charaletris Part-I

 $R = 8.314 \, J \, mol^{-1} \, K^{-1}$

$$e^{-\frac{8000}{8314 \times 300}} = e^{-19.52} - \frac{3.33 \times 10^{-9}}{}$$

[G]

At 45°C

$$E_a = 50 \text{ KJ mod}^{-1} = 50000 \text{ J mod}^{-1}$$

 $T = 45^6 \text{C} + 273 = 318 \text{ K}$
 $R = 8.314 \text{ J mod}^{-1} \text{ R}^{-1}$

$$e^{\frac{3000}{4204.318}} = e^{-18.91} = 6.13 \times 10^{-9}$$

Exponential factor at $25^{\circ}\mathrm{C}=1.73\times10^{-9}$ Exponential factor at $35^{\circ}\mathrm{C}=3.33\times10^{-9}$ Exponential factor at $45^{\circ}\mathrm{C}=6.13\times10^{-9}$

Hence exponential factor doubles for every 10°C rise in T

(iii) Prove that for every 10°C rise in of temperature, the factor doubless and so rate Constant also doubles.

φ

٥

Rate is given by Amhenius eq.

 $k = A_e \cdot \frac{E_e}{\hbar t}$

Por a praction A is almost constant

1 swile coloniations of that fill here. Results of part (ii) shows that, for every LOTC rise in T. exponential factor is

Hence rate is doubled for every 10°C rise in T

H₂ and L₁ react to produce Hi. Pollowing data for rate constant at serious temperatures (R) have been collected. 6.414 × 10

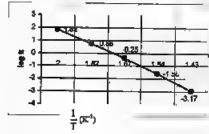




(i) Plot a graph between I/I on x-units and log k on the y-exis.

(ii) Beasure the slope of this straight line and calculate the energy for activation of the

Martin		T was \$	17 👰
500	6.81 x 10⁴	-3 17	2.00 x 10-1
550	2.64 x 10°E	-1:5B	1.82×10 ⁻⁹ ·
600	5.60x10 ⁻¹	-0.25	1.67×104
500	7.31x10°	0.86	1.54°x 10°4
700	6.67x10*1	1,82	I.48 × 10 4



From plotted graph Slope = $-8.36 \times 10^{4} \text{ K}$

 $R = 8.314 \, J \, mol^{-1} \, K^{-1}$

According to Arrhenius equation

$$\log k = \log A - \frac{E_s}{2.303RT}$$

stope =
$$\frac{E_a}{2.309R}$$

or
$$E_a = -2.303 \times R \times slope$$

$$E_a = -2.303 \times 0.314 \times [-8.36 \times 10^{3}]$$

College Characters: Part-1



Multiple Choice Questions from PAST PAPERS

- Specific rate constant is equal to rate of reaction, when concentration of reactants are:
 (Labore board, 2014) (a) Zero
 - (b) Four (c) Three (d) Unity
- 2. The energy of activated complex is manipul tond, 2011 (a), Greater then the reactants & products
 - (b) Less than the reactants & products

 - (c) Equal to the products
 - (d) Equal to the reactants
- If 75% of any given amount of radioactive element district returns in 60 minutes, the half life of radioactive element is; (Separate land, 1819)
 - (a) 20 minutes (b) 30 minutes
- (c) 45 minutes
- (d) 25 minutes
- Sugar solution bydrokyses to glucose and fruction in the presence of ensyme. Get board, 2006: Printed board, 2010) (a) urease (b) invertee (с) зунтаме
- (d) none of these
- When a reaction proceeds in sequence of steps, the overall rate is determined by: (Outcomes board, 2009)
 - (a) fastest step (b) slowest stap
 - (c) molecularity of all steps (d) color of different steps
- 6. Anthenius equation is represented at (Emulsion Board, 1940)
- (a) $A = Ke^{-LAT}$ (b) $K = Ae^{H-Ee}$ (c) $A = Ke^{-H-E}$ (d) $K = Ae^{-AcH}$
- 7 The Hydrolysis of Supar is established by: Makes Brook, steps
 - (n.) Ureane (b) Invertase (c) Zymane (d) Diamines
- (a) Inhibitor (b) retained (c) Promoter (d) Autocataged

 7. The unit of rate constant for zero order reaction is (retained than the first (a) den³ (b) mole dun³ (c) dun³ mole (s² (d) mole of the rate of a character reaction is independent of the material reaction is independent of the mole (s² (d) mole of the rate of a character reaction is independent of the material reaction in the material reaction is independent of the material reaction in the material reaction is independent of the material reaction in the material reaction is independent of the material reaction in the material reaction is independent of the material reaction in the material reaction is independent of the material reaction in the material reaction is independent of the material reaction in the material reaction is independent or reaction in the material reaction in the material reaction is independent or reaction in the material reaction in the 8. A substance which makes the catelyst more effective is called: and

- - [a] Molecularity (b) Temperature (c) nature of reactants
- (c) nature of reception (C) is 5760-years, 100 reg of attempts of Co⁴⁴ will reduce to 25 mg in Recombined Sound, 1910) (b) 2880 years (c) 576500 years (d) 5760 years
- 12. The equation K=As and is called the Road Speed Spe
 - (a) Rate law (4)

(c) Apheneous equation



665 Busties Kings 1.3 A substance which make the catalyst more effective is called (Letow Board, 201). (a) promoter (b) Inhibitor (c) Retarder (d) Poison 14. The order of reaction for the reaction NO+O₃ ---- NO₂+O₂ is (Felenished Board, 2011) fair two (b) three (c) one (d) zero 15. The maintenature associate of energy required for an effective collision is called (Labora Book) (a) Activation energy (b) internal energy (c) Translational energy (d) None of these 16. With increase of 10°C temperature, the rate of reaction becomes double. This increase in rate of reactions due to restanded found, 2007) (a) decrease in the activation energy of reaction (b) decrease in number of collision between the molecules (c) increase in activation-energy of reactants (d) increase in number of effective collisions 17 Normal human body temperature is (Lehers Board, 2021). (a) 37°C (b) 98.6°C (c) 37°C (d) 273 K 18. The unit of rate constant depends on (8-godin Board, 2010) (a) order of reaction (b) molecularity (c) number of reactants all of above The rate of reaction determined at any given time is called (Surgodin Board, 2013) (a) instantaneous rate (b) average rate (c) both 20 Velocity constant is the rate of reaction when the concentrations of the reactants are m) 2910 (b) anily (c) two

Answers to Multiple Choice Questions from Post Popers.								
4na	Q#	Ans	Q#	Ans	Q#	Ans	Q#	Ans
(d)	2	(a)	3	(b)	4	· (b)	5	(b)_
(d)	7	(b)	8	(c)	9		10	(a)
(a)	12	(c)	13	(a)	14		15	(8)
(d)	17	(a)	1B	(a)	19	(a)	20	[6]
֡	Ans (d) (d) (a)	Ans Q# (d) 2 (d) 7 (a) 12	Ans Q# Ans (d) 2 (a) (d) 7 (b) (a) 12 (c)	Ans Q# Ans Q# (d) 2 (a) 3 (d) 7 (b) 8 (a) 12 (c) 13	Ans Q# Ans Q# Ans (d) 2 (a) 3 (b) (d) 7 (b) 8 (c) (a) 12 (c) 13 (a)	Ans Q# Ans Q# Ans Q# (d) 2 (e) 3 (b) 4 (d) 7 (b) 8 (c) 9 (e) 12 (c) 13 (a) 14	Ans Q# Ans Q# Ans Q# Ans (d) 2 (e) 3 (b) 4 (b) (d) 7 (b) 8 (c) 9 (b) (a) 12 (c) 13 (a) 14 (a)	Ans Q# Ans Q# Ans Q# Ans Q# (d) 2 (a) 3 (b) 4 (b) 5 (d) 7 (b) 8 (c) 9 (b) 10 (a) 12 (c) 13 (a) 14 (a) 15

Detailed Explanation of Past Papers MCQs & Answers to all Past Papers SHORT QUESTONS in COLLEGE CHEMISTRY OBJECTIVE BOOK! o Chambers Post-I 659

SHORT & LONG QUESTIONS FROM PAST CHEMICAL KINETICS, RATE OF REACTION

(1) What do you mean by chemical kinetics? (0-

(2) What is meant by rate of reaction? Describe graphically, glass to

(3) Define rate of chemical reaction and give its units. (Laters Sound, 2012) (4) Define reaction iduatics and rate of reaction pain

(5) Define instantaneous and average rate of reaction, and seed, with

(5) Differentiate between average and instantaneous rise of reaction, a. Habanatur Board, 2009. Surpodus Board, 2010, 2013, 2014: Rendpint Board. 2013) Define specific rate constant. (Labora Board, 2013)

(7) Differentiate between rate and rate constant of a reaction granula Balancelour Bound, 2004, 2011)

(1) Define (1) Rate of reaction (11) Specific rate of a reaction (18) Order of a reaction. Baselystedi Board, 2009: Falsahdad Board, 2011)

ORDER OF REACTION

Short Omentions

(1, Define order of reaction with examples, flater Bend, 2011, 2014; Amil Kin

(5) How the mechanism of a chemical reaction can help to point out the rate of determining Step? (Emolyind Sound, 2007)

Luna Circuitors

(1) Explain with one example first order, second order, third order and pseudo first order

reaction: (Behvelper Board, 1018) ENERGY OF ACTIVATION

(1) Define energy of activation. What is the effect of semperature on the activation energy of

O reaction. Francisched Samt. 2009. p.G. Rom Sord, 2019.

Deffine (a) Activation energy. (b) Activated complet Robus South, 2008. Submidger Samt. 2010. Francische Samt. 2010. Francische Samt. 2011. On Under what conditions on callected complex is formed (Schamolger Board, 2011).

(Schamolger Board, 2011).

What is a name of a characters? What is its significance? (Samtyland Samt., 2019).

(3) What is energy, of activation? What is to significance? (Company Dec.)

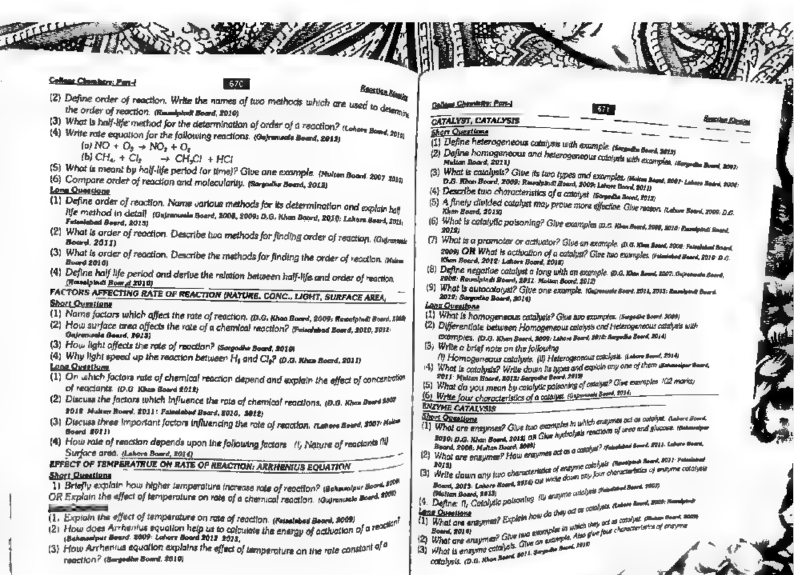
Lena Charatters

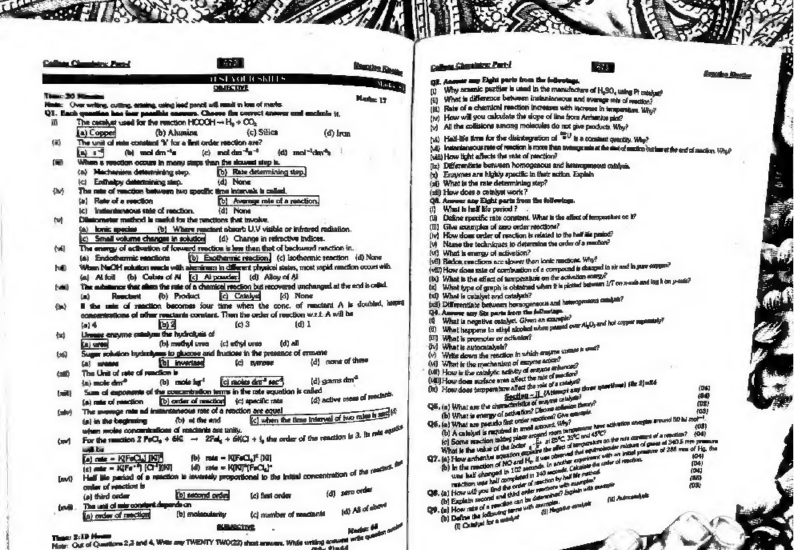
[1] Write a brief note on energy of activation. Explain such propins that Maham Bound, 2019: Maham Bound, 2019: Calmand Bound, 2019: Calmand Bound, 2019: Calmand Bound, 2019: Maham Bound, 2019: Calmand Bound, 2019: Calmand Bound, 2019: Calmand Bound, 2019: Calmand, 2019: Ca

(3) What is collision theory of chamical received personal sections (3)

DETERMINATION OF ORDER OF REACHOR, RATE DESIGNATION STEP Short OverHank

(1) Write names of any faut methods for finding out order of reaction.





674 MODEL PAPER
PUNJAB BOARDS OF INTERMEDIATE AND SECONDARY EDUCATION

Paper Code

Time Allowed: 20 minnies

Mote:- You have four choices for each objective type question as A, B, C and D. The choice which you think is correct: till that circle in frust of that question number. Use marker or you to fill the circles. Custing or filling two or more circles will trouble in zero mark in that question.

Marie and	Quantion	A	is.	C	
1	Empirical formula of Giucoses	CHO	CHO	CHO,	D
-	The number of molecules present to 9 (6.02 × 10 ²³	9.03 × 10°	C,H,O 1.20 x 10
	The drying agent used in a designative is	Lithium	Sodium Chloride	Potaminth Chlorida	Calchem
	The highest temperature at which a solution con exert as liquid, is called it	Absointe	Consolute	Critical	Chioride Transition
	The boding point of water at Mount Everest is	69oC	74°C	79°C	Temperatu B4 ⁹ C
	The existence of an element in more than one crystalline forms is known as	hotropy	Aniosotropy	Enthopy	Allotropy
7	The Scientist Charlesick in 1932 discovered	Proton	Neutron.	Electron	Position
	The values of Quantum numbers for 3P orbital are	n = 1, i = 1	n = 2 . i = I	n = 3 , e = 1	n = 3 , ! = 1
•	The compound which follows justice rule for bonding is	NeC	BCI,	FF;	SF.
10	The Highest percentage of lonic character is in	HF	на	HBr	HÜ
		Enthalpy of Formation	Enthalpy of atomisation	Enthalpy of reaction	Enthalpy of combustion
	In Haber's process, the mercimum yield of summonia can be obtained by	Increasing Pressure	Decreasing	ncreesing volume	increasing emperature
	The selt desolved in seeler forms a solution with pH greater than 7 is	NaCl	Na ₂ CO ₂	CuSO ₄	NHA
	The elevation of boding point of 0.1 molal solution is	0.0052°C	0.062°C	0.52°C	5.2 ⁶ C
1	OF, in	+1	-1	+ 2	-2
	In Lead Accumulator call, the electrolytes	20 % H,SO,	30 % H ₂ SO ₄	40 % H ₂ SO ₄	50 % H ₂ 50 ₄
	Sucrose is consented into Clucose & inuctose by enzyme catalyst called	enverinee	Maltage	(Jenne	J 150

575

Model Paper Chamistry Subjection

ficturoredinta Part - I (11th Class) Examination Section 2015-2016 as Total marks: 55 Time: 3:10 hours SECTION ____

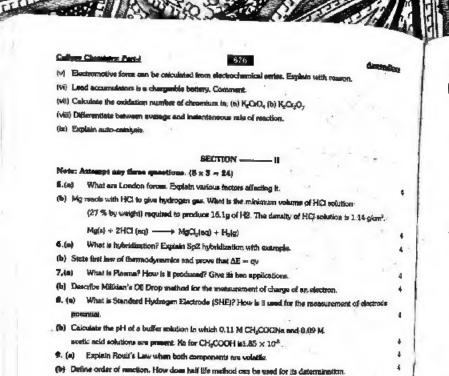
2. Asswer any Eight parts from the followings:

- The removal of an electron from a neutral atom is an endothernic process. English with numero. 8 × 2 × 14
- (ii) Actual yield is always less than theoretical yield. Give two resent.
- (iii) Calculate the no. of molecules present in 34 g of H,PO,
- (iv) Solvent extraction ferms the Distribution Law Justify.
- (v) Define sublimation. Give one example.
- (vi) Calculate the value of General Gas constant in Sturits.
- (vii) Pilote feel uncomfortable breathing at higher atritide. Give reason
- (viii) Genes deviate from ideal behaviour at low respensive and high pressure. Give re-
- (bt) Table soft is an insulator in solid state, Justily.
- (x) Liquid crystals can be used in diagonous of Canter Explan.
- (Al) Evaporation is a cooling process. Give remon.
- (xii) Graphite has dippery touch: Give resear.

8 = 2 = 16

- S. Answer any Eight parts from the followings:
- (i) Positive rate are also called carel tops. One reserv. (ii) The radius of first orbit of hydrogen atom is 0.569 A^o. Calculate the radius of Srd orbit of hydrogen atom
- (iii) Explain mark effect.
- (iv) Paramere can effect the production of Cathoda Reys.
- (v) Dipole interment of CO_p is zero. While that of FE₀ is 1.65 D. Explain.
- (vi) Explain the geometry of H₂Se molecule.
- (vii) Electronegativity incremes from hell to right in puriodic table. Give sem
- (viii) Sketch the molecular orbital picture of \mathcal{O}_{y}
- (its) Enthalpy is a state function. Austly.
- (x) Born Huber's Cycle is another form of Ham's Law, Justily. 04) Bulliers are imposted in many areast of Chemistry, Junely

- (xii) Define Le-Chateller's principle
- 4. Answer any fits parts from the fallowing. (i) Give the applications of the solutility product.
- (ii) Depression of freezing point is a collegative property. Audit.
- [iii] Ne₂5O₄ , 10H₂O group discontinuous schillis, care Che reserv (av) What is the modeliny of a solution prepared by discourse 5 g of Obscore in 250g of



			dagment	mits	, осними
Length	meter	m	Valuing	åtre	1 L (10 m)
Mass	kilogram	lig .	Length Z	angitirom	Aº (0.1nm)
Time	second	1	Pressure K	atmosphere toer	stm(101.325kPa) fromHg(133.32Pa)
Temperature	Helvin	K	Energy 🐉	calionie electron volt	eV (1.602×10 PJ)
Electrical 🕌	ompere	апар	Lempiroture	degree celuus	°C (K-273, 15)
Luquinous intensity	candela	cd	Croprophotion	molecity	M (mol/L or mol/des ²)
Amount of \$	mole	mol	100		

6 2	Fable C-2 trac Fraction	ar d Maltipiles for fise SI	
113.00.5	1031	duci. d	104
csa. E	1011	centi. S	10*
peta. P		milli, en	104
era, T	10**	micro. Its	104
giga: G	10		10-9
niega, M.	10*	graffic. #	10°03
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rcto. tr	10	tentig. I	10-m

Lode	e (3 Cremon Berned Lan	Sunfed -
	Name of with	1 (ing-m / r)
Physical Quantity	out	Hz (cycles / s)
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	alt	C (amp-s)
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the same of charge	H	O.P./mp
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		F(C/V)
petrical Constitute	41	1

